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2.	Patent application number (The Patent Office will fill in this part)	200258.2	07 JAN 2002	
	Full name, address and postcode of the or of each applicant (underline all surnames)	NORCHIP AS INDUSTRIVEIEN 8 N-3490 KLOKKARS NORWAY	TUA	
	Patents ADP number (if you know it) If the applicant is a corporate body, give the country/state of its incorporation	NORWAY	794673	
4.	Title of the invention	DETECTION OF E	IUMAN PAPILLOMA	AVIRUS E6 mRNA
5.	Name of your agent (if you have one) "Address for service" in the United Kingdom to which all correspondence should be sent (including the postcode)	BOULT WADE TO VERULAM GARI 70 GRAY'S INN R LONDON WC1X	DENS OAD	
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7.	If this application is divided or otherwise derived from an earlier UK application, give the number and the filing date of the earlier application	i Number of earlier	application	Date of filing (day / month / year)
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11		I/We request the grant of a patent on the b	pasis of this application.
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DETECTION OF HUMAN PAPILLOMAVIRUS E6 mRNA

The present invention is concerned with oligonucleotide primers and probes for use in detecting the presence of mRNA transcripts from the E6 gene of human papillomavirus in clinical samples.

In the last few years, there has been an improvement in the methods used to detect HPV, with methods based on amplification of nucleic acids using the polymerase chain reaction (PCR) becoming 10 increasingly widespread. It is now possible to detect small amounts of HPV DNA (<100 pg), quantify the amount of viral DNA in clinical samples, identify a broad spectrum of genital HPV types, test for selected HPV types and localise the viral genome transcripts 15 and proteins to the individual cells. Since HPV detection is often carried out in the presence of vast quantities of host nucleic acids and cells not infected with the virus, the ability of the primers to be virus specific is critical for a sensitive and 20 specific amplification.

The present inventors have selected new primer and probe sequences, specific for the E6 region, which may be used in the detection of E6 transcripts by the NASBA technique, particularly sensitive, real-time NASBA, or by RT-PCR. The inventors' approach is based upon the development of primers specific for regions of E6 which are conserved across high-risk, cancer-associated HPV types.

Therefore, in accordance with a first aspect the invention provides target-specific primers and oligonucleotide probes for use in the detection of human papillomavirus (HPV) E6 mRNA, particularly for use in detection of HPV E6 mRNA by RT-PCR or NASBA.

In particular, the invention provides primer and probe oligonucleotides comprising the HPV-specific sequences represented as sequence numbers (SEQ NO) 1 to 133 in Table 1. For each individual sequence an indication is given in the column "primer/probe type" of the general types of primers or probes into which the HPV-specific sequence may be incorporated for the purposes of HPV detection. The HPV type and position in the HPV genome is also indicated.

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Table 1-Summary of primer sequences

г	PRIMER/PROBE	SEQUENCE	SEQ	HPV	nt
ļ	TYPE	SHQUINGE	NO		
15	NASBA P2/PCR	CCACAGGAGCGACCCAGAAAGTTA	1	16	116
13	NASBA P1/PCR	ACGGTTTGTTGTATTGCTGTTC	2	16	368
}	NASBA P2/PCR	CCACAGGAGCGACCCAGAAA	3	16	116
ł	NASBA P1/PCR	GGTTTGTTGTATTGCTGTTC	4	16	368
•	NASBA P1/PCR	TCACGTCGCAGTAACTGT	126	16	208
20	NASBA P1/PCR	TTGCTTGCAGTACACACA	127	16	191
20	NASBA P1/PCR	TGCAGTACACACATTCTA	128	16	186
	NASBA P1/PCR	GCAGTACACACATTCTAA	129	16	185
	NASBA P2/PCR	ACAGTTATGCACAGAGCT	130	16	142
	PROBE				
25	NASBA P2/PCR	ATATTAGAATGTGTGTAC	131	16	182
23			1		
	PROBE NASBA P2/PCR	TTAGAATGŢGTGTACTGC	132	16	185
		TIAMETOJOTOTI			
	PROBE PO (PCP	AATGTGTGTACTGCAAG	133	16	188
	NASBA P2/PCR	AAIGIGIGIACIGCAIC			
30	PROBE	CTTTGCTTTTCGGGATTTATGC	5	16	235
	PROBE	TATGACTTTGCTTTTCGGGA	6	16	230
	PROBE	CAGAGGAGGAGGATGAAATAGTA	7	16	656
	NASBA P2/PCR	GCACAACCGAAGCGTAGAGTCACAC	8	16	741
	NASBA P1/PCR	TGGACAACCGAACCGGACAGACC	9	16	687
35	PROBE	CAGAGGAGGAGGATGAAATAGA	10	16	656
	NASBA P2/PCR	GCACAACCGAAGCGTAGAGTCA	111	16	741
	NASBA P1/PCR	AGCAGAACCGACAGAGCCCATTA	12	16	693
	PROBE	ACGATGAAATAGATGGAGTT	13	18	702
	NASBA P2/PCR	CACGGACACACAAAGGACAG \	14	18	869
40	NASBA P1/PCR	AGCCGAACCACAACGTCACA	15	18	748
	PROBE	GAAAACGATGAAATAGATGGAG	16	18	698
	NASBA P2/PCR	ACACCACGGACACACAAAGGACAG	17	18	869
	NASBA P1/PCR	GAACCACAGGACACACAAAGGACAC	18	18	752
	PROBE	TTCCGGTTGACCTTCTATGT	19	18	651
45	NASBA P2/PCR	GGTCGTCTGCTGAGCTTTCT	20	18	817
	NASBA P1/PCR	GCAAGACATAGAAATAACCTG	21	18	179
	NASBA P2/PCR	GCAAGACATAGAAATAACCTG			

	•	· .	122	18	379	
		ACCCAGTGTTAGTTAGTT	23	18	20-	7
Γ	NASBA P1/PCR	TGCAAGACAGTATTGGAACT		31	16	4
Ī	PRODU	GGAAATACCCTACGATGAAC	24	31	42	3
Ţ	NASDA LZ/	CONCANCIGETCTTTGACA	25	31	26	
	NASBA P1/PCR	ATAGGACGACACACACGGAG	26	31	16	
5	PROBE	GGAAATACCCTACGATGAACTA	27	31	42	
	NASBA P2/PCR	CTGGACACAACGGTCTTTGACA	28	31	126	
	NASBA P1/PCR	TAGGGACACACCACAGGA	29	31	63	
	PROBE	ACTGACCTCCACTGTTATGA	30	$\frac{31}{31}$		56
	NASBA P2/PCR	TATCTACTTGTGTGCTCTGT	31	$\frac{31}{31}$		37
10	NASBA P1/PCR	GACAAGCAGAACCGGACACATC	32	$\frac{31}{31}$		19
	PROBE	TGACCTCCACTGTTATGAGCAATT	33			66
	NASBA P2/PCR	TGACCTCCACTGTTGTGTGCTCTGT TGCGAATATCTACTTGTGTGTGCTCTA	34	31		86
	NASBA P1/PCR	GGACAAGCAGAACCGGACACCCAA	35	31		17
	PROBE	GGACAAGCAGAACCGG	36	31		09
15	NASBA P2/PCR	ACTGACCTCCACTGTTAT CACGATTCCAAATGAGCCCAT	37	31		518
10	NASBA P1/PCR	CACGATTCCAAATGACCCTAT	38	33		763
	NASBA P2/PCR	TATCCTGAACCAACTGACTG	39	3:		594
	NASBA P1/PCR	TTGACACATAAACGAACTG	40	3		620
	PROBE	CAGATGGACAAGCACCTAT	41	3		807
20	NASBA P2/PCR	TCCTGAACCAACTGACCTAT	42	3		699
20	NASBA P1/PCR	CCCATAAGTAGTTGCTGTAT	43			431
	PROBE	CCACAAGCACCACC	44		13	618
	NASBA P2/PCR	GACCTTTGTGTCCTCAAGAA	45		33	
	NASBA P1/PCR	T ACCTO AGT TOGET TO THE	46	· -	33	543
25	PROBE	I ACAACTGCACTGTGTG	47		35	217
25	NASBA P2/PCF	ATTACAGCGGAGTGAGGTAT	48		35	442
	NASBA P1/PCF		4:		35	655
	NASBA P2/PC	T TO A C. A C	5	0	35	844
	NASBA P1/PC	- 1 Chillian Title Total	. 5	1	35	610
20	NASBA P2/PC	o I CCCTAIRIS AGO I OIL	5	2.	35	770
30	NASBA P1/PC	D CTCAATGIGIGIG	5	3	35	270
	PROBE		5	4	35	692
	PROBE	GACAAGCAAAACCAGACACCTCCAA	- 5	55	35	692
	PROBE	GACAAGCAAAACCAGACACC		6	52	144
٥٢	NASBA P2/PC	CR TTGTGTGAGGTGCTGGAAGAAT		57	52 .	358
35	NASBA P1/PC	CCCTCTCTTCTAATGTTT		58	52	296
				59	52	296
	PROBE NASBA P2/P0	- CMCCCTACGCTTTTTATCIA		60	52	507
	NASBA P1/P	CD CGGGTCTCCAACACTCTGAACA		61	52	461
	PROBE	TCCAAACAAGCGATTTCA		62	58	157
40	NASBA P2/P	CR TCAGGCGTTGGAGACATC		63	58	301
	NASBA P1/P	ACCAATCGTAAGCACACT		64°	58	173
	NASBA P2/P	TCTGTGCATGAAATCGAA		65	58	291
	NASBA P1/E	ACCACACTTTACATACTG		66	58	192
		TO A A TICCGTTGAATGCA		67	58	218
45	PROBE	TTGCAGCGATCTGAGGTATAIG		68	В	514
	PROBE NASBA P2/	TACACTGCTGGACAACAT		69	В	619
	NASBA PZ/	TO THE TOTAL CONTROL OF THE TO		70	В	514
	NASBA P1/	TO CA CACACACATGUA		71	В	69:
	NASBA P2/	CTCACACACAGCAACAGGICA		72	В	59
50	NASBA P1/			73	В	59
	PROBE			74	$-\frac{1}{B}$	69
	PROBE	PCR TGACCTGTTGCTGTGGATGTGA		1 12		

NASBA P1/PCR	_			75	В	832
NASBA P1/PCR CATGCCATAGATTATAGA 77	Ĺ	NASBA P1/PCR	TACCTGAATCGTCCGCCAT	75		
NASBA P1/PCR CACGGCAGGCACTTATTAA 78 C 408		PROBE				
Section	Ī	NASBA P2/PCR				
NASBA P2/PCR GCAGACGACCATACAGCARA So 39 210	ſ	NASBA P1/PCR				
NASBA P1/PCR ACACCGAGTATATA 31 39 344	5	PROBE	AGAATTAGAGAATTAAGA			
NASBA P1/PCR ATAGGACGGGAACCAT 82 39 273	Ī	NASBA P2/PCR	GCAGACGACCACTACAGCAAA			
NASBA P2/PCR TATGCTGGACTCGTTT	Ì	NASBA P1/PCR	ACACCGAGTCCGAGTAATA			
NASBA P1/PCR CTTGGGTTTCTTCTGTGTTA	Ì	PROBE	ATAGGGACGGGAACCACT			
NASBA P1/PCR CTTGGGTTTCTCTGTGTA	Ť	NASBA P2/PCR	TATTACTCGGACTCGGTGT	83		
PROBE GGACACAAAACGGGAGGAC 85 39 703 NASBA P1/PCR GCACACCAGGACACAAA 87 39 886 PROBE TAGCCAGCGGACACACAA 87 39 886 PROBE TAGCCAGCGGACACACAA 87 39 886 PROBE TAGCCAGCGGACACACAC 88 39 749 NASBA P2/PCR AACCATTGAACCACGGAA 89 45 430 NASBA P1/PCR TCTTCTTGCCTGCTGCTA 90 45 527 NASBA P1/PCR TAGCTATCACCAGCAGAAA 91 45 428 NASBA P1/PCR TAGCTATCATCTGGTTTCCCTACG 92 45 558 PROBE GTACCGAGGAGATATA 93 45 467 PROBE GTACCGAGGAGATATCACA 94 45 467 NASBA P1/PCR TAGCTATCACTGGTTTCCCTACG 92 45 566 NASBA P1/PCR CTGTACTGTGTTTACCA 94 45 467 NASBA P1/PCR CTGTTGACTGTTGTTTACCA 95 45 666 NASBA P1/PCR CACCACGACACAAAGGACAAG 96 45 868 NASBA P1/PCR CACCACGACCACAAAGGACAAG 98 45 868 NASBA P1/PCR CACCACGACCACAAAGGACAAG 98 45 868 NASBA P1/PCR CACGACACACAAAGGACAAG 98 45 868 PROBE GAGTCACAGAGGACAAA 100 45 866 PROBE AGGAAACCATCACAAGGACAAG 100 45 866 PROBE AGGAAACCACAAAGGACAAG 100 45 666 PROBE AGGAAACGATGAAGACATG 101 45 686 PROBE AGGAAACGATGAAGACATG 101 45 686 PROBE AGGAAACATCCCGCCGAGGACCAAA 103 45 730 NASBA P1/PCR CCCCTTTACATCTGCTGT 105 51 807 NASBA P1/PCR ACGGGCAAACCAGATGATA 106 51 655 NASBA P1/PCR ACGGGCAAACCAGGACATAGATA 106 51 655 NASBA P1/PCR ACGGGCAAACCAGGACAAACCAT 107 51 829 PROBE GCAGGTGTTCAACTGTGTAT 107 51 829 PROBE GCAGGTTTCAACTGTTGTTAGA 107 51 829 PROBE GCAGGTTTCAACTGTTGTTAGA 106 51 655 NASBA P1/PCR ACGGGCAAACCAACAACATT 110 56 519 NASBA P1/PCR TTCATCCTCATCTCTCTGA 111 56 665 NASBA P1/PCR CATCCTCATCCTCATCCTCTGA 113 56 665 PROBE AGAGTACACACAACAACATT 114 56 519 NASBA P1/PCR CATCCTCATCCTCAACAACAT 114 56 519 NASBA P1/PCR CATCCTCATCCTCAACAACAT 114 56 519 NASBA P1/PCR CATCCTCATCCTCAACAACATT 114 56 519 NASBA	10		CTTGGGTTTCTCTTCGTGTTA	84		
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PROBE	Ì		GCACACCACGGACACAAA	87	39	886
NASBA P2/PCR AACCATTGAACCCAGCAGAAA 89 45 430					39	749
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NASBA P1/PCR CCACGGACACAAAGGACAAG 98 45 868	ŀ			97	45	654
NASBA P2/PCR GTTGACCTGTTGTTTACGA 99 45 656 NASBA P1/PCR ACGGACACAAAGGACAAG 100 45 868 PROBE GAGTCAGAGGAGAGAGG 101 45 686 PROBE AGGAAAACGATGAGGATG 101 45 696 PROBE AGGAAAACGATGAAGCAGTG 102 45 696 PROBE ACAACTACCAGCCCGACGAGCCGAA 103 45 730 NASBA P2/PCR GGAGGAGGATGAAGTA 104 51 658 NASBA P1/PCR GCCCATTAACATCTGCTGTA 105 51 807 NASBA P1/PCR AGAGGAGGAGGATGAAGTA 106 51 655 NASBA P1/PCR ACGGCAAACCAGGCTTAGT 107 51 829 PROBE GCAGGTGTTCAAGTGTAGT 108 51 747 NASBA P1/PCR ACGGCAAACCAGGCTTAGT 109 51 771 NASBA P2/PCR TTGGGTGCTGGAGACAACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TTGGGTGCTGGAGACAACATC 112 56 520 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 114 56 519 NASBA P1/PCR CATCCTCATCCTCTGA 115 56 764 PROBE AAAGTACCAACGCTGCAAACAT 114 56 519 NASBA P1/PCR CCACAAACTTACACTCACACA 115 56 764 PROBE AAAGTACCAACGCTGCAAGACT 116 56 581 PROBE AAAGTACCAACGCTGCAAGACT 116 56 581 PROBE AAAGTACCAACGCTGCAAGACGT 116 56 581 PROBE AGAACTAACACCTCAAACAGAAAT 117 56 610 PROBE AGAACTAACACCTCAAACAGAAT 117 56 610 PROBE AGAACTAACACCTCAAACAGAGAAT 117 56 610 PROBE AGAACTACACCTCAAACAGAGAGT 118 56 583 PROBE TTGGACAGCTCAAGACGTT 118 56 583 PROBE AGAACTACCACCTTAATT 121 56 410 PROBE GACTATTCATTGAGTGTG 120 56 279 NASBA P1/PCR GACTATTCATTGAGTGTG 122 56 348 PROBE CAACTGACCTTATT 121 56 410 PROBE GACTATTCAGTGTATGAGCC 122 56 348 PROBE CAACTGACCTTATTCATTCATTATTATAGATTATAGATTATAGATTATAGATTATAGATTATAGATTATAGATTATAGATTATAGATTATAGATTATAGAT	}			98	45	868
NASBA P1/PCR ACGGACACAAAGGACAAG 100 45 868	25			99	45	656
PROBE GAGTCAGAGGAGAAAACGATG 101 45 686 PROBE AGGAAAACGATGAGAGTGAGT 102 45 696 PROBE ACAACTACCAGCCCGACGAGCCGAA 103 45 730 RASBA P2/PCR GGAGGAGGATGAAGTAGATA 104 51 658 NASBA P1/PCR GCCCATTAACATCTGCTGTA 105 51 807 NASBA P2/PCR ACAGGAGGAGGAGATAGATA 106 51 655 NASBA P1/PCR ACAGGAGGAGAGTAGATA 106 51 655 NASBA P1/PCR ACAGGAGAGAGTAGATA 106 51 655 NASBA P1/PCR ACAGGAGAGAGTAGATA 108 51 747 PROBE GCAGTGTTCAAGTGTAGT 108 51 747 ASBA P2/PCR TTGGGATGGAGACAACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TTGGGGTGCTGGAGACAACATC 112 56 520 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR TTGGGGTGCTGGAGACAACATC 112 56 520 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCAGACA 115 56 519 NASBA P1/PCR CATCCTCATCCTCAGACA 115 56 764 PROBE AAAGTACCACGCTGCAAGACAT 116 56 581 PROBE ACAACTACACGCTGCAAGACAT 117 56 610 PROBE ACAACTACACCTCAACAGAGAAAT 117 56 610 PROBE ACTACCACGCTGCAAGACGT 118 56 583 ASBA P2/PCR GACTATTCCTTATGCAGTCTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACTATTCAGTGTATGAGC 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A PROBE CAACTGAYCTMYACTGTTATGA 124 A PROB	2.5			100	45	868
PROBE AGGAAACGATGAAGCAGATGAGT 102 45 696 PROBE ACAACTACCAGCCCGACGAGCCGAA 103 45 730 NASBA P2/PCR GGAGGAGGATGAAGTAACATA 104 51 658 NASBA P1/PCR GCCCATTAACATCTGCTGTA 105 51 807 NASBA P2/PCR AGAGGAGGATGAAGTAGATA 106 51 655 NASBA P1/PCR ACGGGCCAGGAGCTAGATA 106 51 655 NASBA P1/PCR ACGGGCAAACCAGGCTTAGT 107 51 829 PROBE GCAGGTGTTCAAGTGTAGT 108 51 747 35 PROBE TGGCAGTGGAAAGCAGGTGGAGACA 109 51 771 NASBA P2/PCR TTGGGGTGCTGGAGACAACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR CATCCTCATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCATCCTCTGA 114 56 519 NASBA P1/PCR CATCCTCATCCTCATCCTCTGA 115 56 764 PROBE AAAGTACCAACGCTGCAGACAT 116 56 581 PROBE AGAACTACACCTCAACACA 115 56 764 PROBE AGAACTACACCTCAACACA 115 56 610 PROBE AGAACTACACCTCAACAGAAAT 117 56 610 PROBE AGAACTACACCTCAACAGAGGT 118 56 583 NASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGGAGCCTTATT 121 56 410 PROBE GACTATTCAGTGTATGGAGC 122 56 348 PROBE GACTATTCAGTGTATGAGCC 122 56 348 PROBE CAACTGGACCTTATTATGA 123 A PROBE CAACTGGACCTTAYCTGTTATGA 124 A	1			101	45	686
PROBE ACAACTACCAGCCCGACGAGCCGAA 103 45 730 NASBA P2/PCR GGAGGAGGATGAAGTAA 104 51 658 NASBA P1/PCR GCCCATTAACATCTGCTGTA 105 51 807 NASBA P2/PCR AGAGGAGGAGGATGAAGTAA 106 51 655 NASBA P1/PCR ACGGGCAAACCAGGCTTAGT 107 51 829 PROBE GCAGGTGTTCAAGTGTAGTA 108 51 747 PROBE TGGCAGTGGAAACCAGGAGCAA 109 51 771 NASBA P2/PCR TTGGGGTGCTGGAGACAACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TTGATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCATCCTCTGA 114 56 519 NASBA P1/PCR CATCCTCATCACACCACACA 115 56 564 PROBE AAAGTACCAACGCTGCAACACA 115 56 764 PROBE AGAACTAACACCTCAACACA 116 56 581 PROBE AGAACTAACACCTCAAACAACAT 117 56 610 PROBE AGAACTAACACCTCAAACAGAAAT 117 56 610 PROBE AGAACTAACACCTCAACAGACGTT 118 56 583 PROBE ATTGGACGCTGCAAGAGCGTT 118 56 583 ASSA P1/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAACACCTTATT 121 56 410 PROBE GACATCTGTAACACCTTAATT 121 56 410 PROBE GACATCTGTAACACCTTAATT 121 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A PROBE CAACTGAYCTMYACTGTTATGA 124 A TATTATATA				102	45	696
NASBA P2/PCR GGAGGAGGATGAAGTAGATA 104 51 658 NASBA P1/PCR GCCCATTAACATCTGCTGTA 105 51 807 NASBA P2/PCR AGAGGAGGAGGATGAAGTAGATA 106 51 655 NASBA P1/PCR ACGGGCAAACCAGGCTTAGT 107 51 829 PROBE GCAGGTGTTCAAGTGTAGTA 108 51 747 PROBE TGGCAGTGGAAAGCAGTGGAACA 109 51 771 NASBA P2/PCR TTGGGGTGCTGGAGACAACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TTGGGGTGCTGGAGACAAACATC 112 56 520 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 114 56 519 NASBA P1/PCR CCACAAACTTACACTCACACA 115 56 764 PROBE AAAGTACCACCTCAACACA 115 56 581 PROBE AGAACTAACACCTCAACACA 117 56 610 PROBE AGAACTAACACCTCAACACGTT 118 56 583 ASBA P1/PCR GACATCTGTAGAGGGT 119 56 656 NASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACATCTGTAGCACCTTATT 121 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A PROBE CAACTGAYCTMYACTGTTATGA 124 A SO PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A SO PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A				103	45	730
NASBA P1/PCR GCCCATTAACATCTGCTGTA 105 51 807 NASBA P1/PCR AGAGGAGGAGGATGAAGTA 106 51 655 NASBA P1/PCR AGAGGAGGAGGATGAAGTA 106 51 655 NASBA P1/PCR ACGGGCAAACCAGGCTTAGT 107 51 829 PROBE GCAGGTGTTCAAGTGTAGTA 108 51 747 PROBE TGGCAGTGGAAAGCAGTGGAGACA 109 51 771 NASBA P2/PCR TTGGGGTGCTGGAGACAACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P2/PCR TGGGGTGCTGGAGACAACATC 112 56 520 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 114 56 519 NASBA P1/PCR CATCCTCATCCTCTGA 115 56 764 PROBE AAAGTACCAACACACACACACACACACACACACACACACA	20			104	51	658
NASBA P1/PCR ACGGCAGAGAGAGATAAACATA 106 51 655 NASBA P1/PCR ACGGCAAACCAGGCTTAGT 107 51 829 PROBE GCAGGTGTTCAAGTGTAGTA 108 51 747 PROBE TGGCAGTGGAAAGCAGTGGAGACA 109 51 771 NASBA P1/PCR TTGGGGTGCTGGAGACAACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TGGGGTGCTGGAGACAACATC 112 56 520 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 114 56 519 NASBA P1/PCR CCACAAACTTACACTCACACA 115 56 764 PROBE AAAGTACCAACGCTGCAAGACAT 116 56 581 PROBE AGAACTAACACCTCAAACAGAAAT 117 56 610 PROBE AGAACTAACACCTCAAACAGAAAT 117 56 610 PROBE AGAACTAACACCTCAAACAGAAAT 117 56 656 NASBA P1/PCR GACATCTGTAGAGAGGT 118 56 583 45 PROBE TTGGACAGCTCAGAGAGAGT 118 56 583 NASBA P1/PCR GACATCTGTAGCAGCTT 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACATTTCAGTTATGAGG 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A	30				51	807
NASBA P1/PCR ACGGCAAACCAGGCTTAGT 107 51 829					51	655
PROBE GCAGGTGTTCAAGTGTAGTA 108 51 747						829
PROBE TGGCAGTGGAAACATCT 109 51 771 NASBA P2/PCR TTGGGGTGCTGGAGACAACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P2/PCR TGGGGTGCTGGAGACAACATC 112 56 520 NASBA P1/PCR TGGGGTGCTGGAGACAACATC 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCATCCTGA 114 56 519 NASBA P1/PCR CCACAAACTTACACTCACAACA 115 56 764 PROBE AAAGTACCAACCACACACA 115 56 764 PROBE AGAACTAACACCTCAAACAGACT 116 56 581 PROBE AGAACTAACACCTCAAACAGAAAT 117 56 610 PROBE AGTACCAACGTGCAAGACGTT 118 56 583 ASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACTATTCAGTGTAGGAGC 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A SO PROBE GAAMCAACTGACCTATT 124 A SO PROBE GAAMCAACTGACCTATCTATT 125 A SO PROBE GAAMCAACTGACCTATCTATT 124 A SO PROBE GAAMCAACTGACCTATCTATT 125 A SO PROBE GAAMCAACTGACCTATCTATTATTATAGA 125 A SO PROBE GAAMCAACTGACCTATCTATTATTATAGA 125 A SO PROBE GAAMCAACTGACCTATCTATTATTATAGA 125 A SO PROBE GAAMCAACTGACCTATCTATTATTAGA 126 A SO PROBE GAAMCAACTGACCTATCTATTATTAGA 127 A SO PROBE GAAMCAACTGACCTATCTATTATTAGA 126 A SO PROBE GAAMCAACTGACCTATCTATTATTAGA 127 A SO PROBE GAAMCAACTGACCTATATTATTAGA 127 A SO PROBE GAAMCAACTGACCTATTATTAGA 127 A SO PROBE GAAMCAACTGACCTATATTATTATTATTAGA 127 A SO PROBE GAAMCAACTGACCTATATTATTATTATTATTATTATTATTATTATTATTA					51	747
NASBA P2/PCR TTGGGGTGCTGGAGACATCT 110 56 519 NASBA P1/PCR TTCATCCTCATCCTCTGA 111 56 665 NASBA P1/PCR TTGATCCTCATCCTCTGA 111 56 520 NASBA P1/PCR TGGGGTGCTGGAGACAACATC 112 56 520 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P1/PCR CATCCTCATCCTCTGA 114 56 519 NASBA P1/PCR CCACAAACTTACACTCACAACA 115 56 764 PROBE AAAGTACCAACGCTGCAAGACGT 116 56 581 PROBE AGAACTAACACCTCAAACAGAAAT 117 56 610 PROBE AGTACCAACGCTGCAAGACGT 118 56 583 PROBE AGTACCAACGCTGCAAGACGT 118 56 583 PROBE TTGGACAGCTCAGAGACGT 118 56 656 NASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACATCTGTAGCACCTTATGA 123 A PROBE CAACTGAYCTMYACTGTTATGA 124 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A So So So So So So So	25					771
NASBA P2/PCR TTGGGGTGCTGAAACATC 111 56 665 NASBA P1/PCR TTCATCCTCATCCTCTGA 112 56 520 NASBA P2/PCR TGGGGTGCTGGAGACAACATC 112 56 520 NASBA P1/PCR CATCCTCATCCTCTGA 113 56 665 NASBA P2/PCR TTGGGGTGCTGGAGACAACAT 114 56 519 NASBA P1/PCR CCACAAACTTACACTCACAACA 115 56 764 PROBE AAAGTACCAACGCTGCAAGACGT 116 56 581 PROBE AGAACTAACACCTCAAACAGAAAT 117 56 610 PROBE AGTACCAACGCTGCAAGACGTT 118 56 583 PROBE TTGGACAGCTCAGAGAGTT 118 56 583 PROBE TTGGACAGCTCAGAGGATGAGG 119 56 656 NASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACATCTGTATGGAGC 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A Solvent 125 A S	35					519
NASBA P1/PCR						665
NASBA P2/PCR TGGGGTGCTGGAGACAAACATCAACATCAACATCAACATCAACATCAACAACA						520
NASBA P1/PCR						
NASBA P2/PCR	4.0					
PROBE AGACTACACCTCACAGACT 116 56 581 PROBE AGACTAACACCTCACAGACGT 116 56 610 PROBE AGACTAACACCTCACACAGACAT 117 56 610 PROBE AGTACCACCCTCACAGACGTT 118 56 583 PROBE TTGGACAGCTCAGAGGATGAGG 119 56 656 NASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACTATTCAGTGTATGAGC 122 56 348 PROBE CACTGAYCTMYACTGTTATGA 123 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A	40					
PROBE AGAACTAACACCTCAAACAGAAAT 117 56 610 PROBE AGTACCAACGCTGCAAGACGTT 118 56 583 PROBE TTGGACAGCTCAGAGGATGAGG 119 56 656 NASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACTATTCAGTGTATGAGC 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A						
PROBE AGTACCAACGCTGCAAGACGTT 118 56 583 PROBE TTGGACAGCTCAGAGGATGAGG 119 56 656 NASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACTATTCAGTGTATGAGC 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A						
45 PROBE TTGGACAGCTCAGAGGATGAGG 119 56 656 NASBA P2/PCR GATTTTCCTTATGCAGTGTG 120 56 279 NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACTATTCAGTGTATGAGC 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A						
PROBE TIGGACAGCTCAGAGGATGAGGATGAGGATGAGGATGAGGATGAGGATGAGGATGAGGATGAGGATGAGGATGAGGATGAGGAG	, 					
NASBA P2/PCR GATTITCCTTATGCACTGT NASBA P1/PCR GACATCTGTAGCACCTTATT 121 56 410 PROBE GACTATTCAGTGTATGGAGC 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A	45					
PROBE GACTGTATGAGCC 122 56 348 PROBE CAACTGAYCTMYACTGTTATGA 123 A PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 125 A						
PROBE GACTATICAGISTATICA PROBE CAACTGAYCTMYACTGTTATGA 123 A 50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A 135 P						
50 PROBE GAAMCAACTGACCTAYWCTGCTAT 124 A						+
JU PROBE GRANICATION CONC. 125 B						
PROBE AAGACATTATTCAGACTC 123 A	50					
		PROBE	AAGACATTATTCAGACTC	125		1

Oligonucleotides for use as NASBA P1 primers have the general structure " X_1 -SEQ", wherein " X_1 " represents a nucleotide sequence comprising a promoter and "SEQ" represents the HPV-specific sequence, as given in Table 1. The inclusion of a promoter sequence is essential in NASBA P1 primers but is not necessary in PCR primers, as discussed below. In a preferred embodiment, X_1 may be a sequence comprising a bacteriophage promoter, preferably the T7 promoter. In the most preferred embodiment, X_1 represents the sequence AATTCTAATACGACTCACTATAGGGAGAAGG.

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The oligonucleotide molecules of the invention are selected to be specific for mRNA transcribed from the HPV E6 gene. Active expression of the E7 and E6 genes of HPV is associated with cervical cytological 15 abnormalities which often progress to more serious disease. A number of studies relate the expression of the E6 and E7 genes to oncogenesis. Co-operation between E6 and E7 increases significantly the frequency of immortalization. Evidence has been 20 presented that the E6 and E7 open reading frames are involved in the transforming activity of the virus (Tanaka et al., J. Virol. 63: 1465-1469, 1989). transformation effects of E6 and E7 may at least in part be explained by their interaction with the 25 cellular tumour suppressor gene products p53 and pRb 105, respectively (Boyer et al., Cancer Research. 56: 4620-4624, 1996; Lechner et al. EMBO J. 11: 3045-3051, 1992). 30

HPV16 mRNA isolated from transfected cells and a variety of tumour cell lines and lesions containing both extrachromosomal and integrated HPV16 genomes has

been analysed in multiple laboratories (see Doorbar JA et al., Virology 178:254?262, Rohlfs et al., Virology 183:331?342; Sherman et al., Int. J. Cancer 50:356?364). These studies have shown that several different alternatively spliced transcripts may be produced from the E6 and E7 region. In summary, there are four major transcripts: one with the whole E6/E7 gene area (E6), one with a loss of a coding sequence between basepairs 226 and 409 (E6*I), one with a loss of a coding sequence in a larger part of

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with a loss of a coding sequence in a larger part of
E6 between 226 and 526 (E6*II) and one with the loss
of the E7 transcript (E6*III). However there are
clearly consensus sequences in the area up to 226
basepairs in the E6 region. The inventors therefore
selected the areas between 97 and 226 and between 526
and 880 as areas to target for diagnostic purposes.

The oligonucleotides provided by the invention may be grouped according to specificity for different specific HPV types or groups of HPV types. Sequence 20 numbers 1-12 and 126-133 are specific for HPV type 16, sequence numbers 13-23 are specific for HPV type 18, sequence numbers 24-37 are specific for HPV type 31, sequence numbers 38-46 are specific for HPV type 33. HPV types 16, 18 , 31 and 33 are the major 25 cancer-associated HPV types. Sequence numbers 47-55 are specific for HPV type 35, sequence numbers 56-61 are specific for HPV type 52, sequence numbers 62-67 are specific for HPV type 58, sequence numbers 80-88 are specific for HPV type 39, sequence numbers 89-103 30 are specific for HPV type 45, sequence numbers 104-109 are specific for HPV type 51, sequence numbers 110-122 are specific for HPV type 56. Sequence numbers 68-76 are consensus sequences for group B HPV types (in

particular HPV types 6 and 11). Sequence numbers 77-79 and 125 are consensus sequences for group C HPV types (including HPV types 18, 39 and 45). Sequence numbers 123 and 124 are consensus probe sequences for group A HPV types. Sequence 123 is a consensus for HPV types 16, 31 and 35; sequence 124 is a consensus for HPV types 33, 52 and 58).

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The oligonucleotide molecules of the invention are preferably single stranded DNA molecules. Non-natural synthetic polynucleotides which retain the 10 ability to base-pair with a complementary nucleic acid molecule and are also within the scope of the invention, including synthetic oligonucleotides which incorporate modified bases and synthetic oligonucleotides wherein the links between individual 15 nucleosides include bonds other than phosphodiester bonds. The oligonucleotide molecules of the invention may be produced according to techniques well known in the art, such as by chemical synthesis using standard apparatus and protocols for oligonucleotide synthesis. 20

The oligonucleotide molecules provided by the invention will typically be isolated single-stranded polynucleotides of no more than 100 bases in length, more typically less than 55 bases in length. For the avoidance of doubt it is hereby stated that the language "oligonucleotide comprising sequence number n" excludes the naturally occurring full-length HPV genomes.

The invention provides several general types of oligonucleotide primers and probes incorporating the HPV-specific sequences listed in Table 1. Typically,

such oligonucleotides may comprise additional, non-HPV sequences, for example sequences which are required for an amplification reaction or which facilitate detection of the products of the amplification reaction. The HPV-specific part of the oligonucleotide may consist of one of the sequences listed in Table 1 in the absence of any other contiguous HPV sequences. However, it will be appreciated that minor variations may be made to the HPV-specific sequences, for example the addition, deletion or substitution of bases, without affecting the ability of the oligonucleotide to bind to its

target sequence and function as a primer or probe to a

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material extent.

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The first type of oligonucleotides are primer 1 oligonucleotides (also referred to herein as NASBA P1 primers), which are oligonucleotides of generally approximately 50 bases in length, containing an 20 average of about 20 bases at the 3' end that are complementary to a region of the target mRNA. Oligonucleotides suitable for use as NASBA P1 primers are denoted "NASBA P1/PCR" in Table 1. The 5' ends of the P1 primer oligonucleotides (represented herein in 25 general terms as X1) comprise a promoter sequence that is recognized by a specific RNA polymerase. Bacteriophage promoters, for example the T7, T3 and SP6 promoters, are preferred for use in the oligonucleotides of the invention, since they provide 30 advantages of high level transcription which is dependent only on binding of the appropriate RNA polymerase. In a preferred embodiment, the 5' terminal sequence of the P1 primer oligonucleotides may comprise the sequence AATTCTAATACGACTCACTATAGGG or

the sequence AATTCTAATACGACTCACTATAGGGAAGG. These sequences contains a T7 promoter, including the transcription initiation site for T7 RNA polymerase. The HPV-specific sequences denoted in Table 1 as "NASBA P1/PCR" are suitable for use in both NASBA P1 primers and standard PCR primers. When these sequences are used as the basis of NASBA P1 primers they have the general structure X_1 -SEQ, wherein X_1 represents a sequence comprising a promoter and SEQ represents the HPV-specific sequence. The promoter sequence X_1 is essential. However, when the same sequences are used as the basis of standard PCR primers it is not necessary to include X_1 . The phrase "sequence number" as used in the claims is to be interpreted accordingly.

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For the avoidance of doubt, the phrase "a NASBA P1 primer comprising sequence number 1" is to be interpreted as requiring the presence of an X1 sequence 5' to the HPV-specific sequence listed as sequence number 1, whereas the phrase "a PCR primer comprising sequence number 1" refers to any suitable PCR primer comprising the HPV-specific sequence, X1 not being an essential feature of a PCR primer. The phrase "an oligonucleotide primer including sequence number n" is taken to encompass NASBA P1, NASBA P2 and PCR primers.

A second type of oligonucleotide provided by the invention are NASBA primer 2 oligonucleotides (also referred to herein as NASBA P2 primers) which generally comprise a sequence of approximately 20 bases substantially identical to a region of the target mRNA. The oligonucleotide sequences denoted in

Table 1 as "NASBA P2/PCR" are suitable for use in both NASBA P1 primers and standard PCR primers.

Oligonucleotides intended for use as NASBA P2 primers may, in a particular but non-limiting embodiment, further comprise a sequence of nucleotides at the 5' end which is unrelated to the target mRNA but which is capable of hybridising to a generic detection probe. The detection probe will preferably be labelled, for example with a fluorescent,

luminescent or enzymatic label. In one embodiment the detection probe is labelled with a label that permits detection using ECL™ technology, although it will be appreciated that the invention is in no way limited to this particular method of detection. In a preferred embodiment the 5' end of the primer 2 oligonucleotides may comprise the sequence GATGCAAGGTCGCATATGAG. This sequence is capable of hybridising to a generic ECL™ probe commercially available from Organon Teknika having the following structure:

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Ru(bpy)₃²⁺-GAT GCA AGG TCG CAT ATG AG-3'

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In a different embodiment the primer 2 oligonucleotide may incorporate "molecular beacons" technology, which is known in the art and described, for example, in WO 95/13399 by Tyagi and Kramer, Nature Biotechnology. 14: 303-308, 1996, to allow for real-time monitoring of the NASBA reaction.

A third type of oligonucleotide molecules provided by the invention are target-specific probe oligonucleotides (denoted "probe" in Table 1). The probe oligonucleotides generally comprise a sequence of approximately 20-25 bases substantially identical

to a region of the target mRNA, or the complement thereof. The probe oligonucleotides may be used as target-specific hybridisation probes for detection of the products of a NASBA or PCR reaction. In this connection the probe oligonucleotides may be coupled to a solid support, such as paramagnetic beads, to form a capture probe (see below). In a preferred embodiment the 5' end of the probe oligonucleotide may be labelled with biotin. The addition of a biotin label facilitates attachment of the probe to a solid support via a biotin/streptavidin or biotin/avidin linkage.

A fourth type of oligonucleotide molecules provided by the invention are target-specific probes incorporating "molecular beacons" technology which is known in the art and described, for example, by Tyagi and Kramer, Nature Biotechnology. 14: 303-308, 1996 and in WO 95/13399.

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The term "molecular beacons probes" as used herein is taken to mean molecules having the structure:

 $X_2-arm_1-target-arm_2-X_3$

wherein "target" represents a target-specific sequence of nucleotides, " X_2 " and " X_3 " represent a fluorescent moiety and a quencher moiety capable of substantially or completely quenching the fluorescence from the fluorescent moiety when the two are held together in close proximity and " arm_1 " and " arm_2 " represent complementary sequences capable of forming a stem duplex.

The invention provides molecular beacons probes incorporating a target-specific sequence comprising one of sequence numbers 6, 18, 35, 43, 123, 124 or 125.

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Suitable pairs of arm_1 and arm_2 sequences for use with these HPV-specific sequences include, but not exclusively, the following:

	exclusively, the following.
10	For use with sequence number 6:
	CGCATGCATGCG
	CCAGCTAGCTGG
	CACGCGCGTG
	CGATCGCGATCG
15	
	For use with sequence number 18:
	CGCATGCATGCG
	CCGTCGCGACGG
	CGGACCGGTCCG
20	CGATCGCGATCG
	For use with sequence number 35:
	CCGAAGGCCTTCGG
	CCGTCGCGACGG
25	CACGTCGCGACGTG
	CGCAGCGCTGCG
	CGATCGCGATCG
	For use with sequence number 43:
30	CCAAGCGCTTGG
	CCAAGCGCGCTTGG
	CCCAGCGCTGGG
	CCAAAGCGCTTTGG
	CCTGCGCAGG
	= = ··

CGATCG-----CGATCG

For use with sequence number 123:

CGCATG-----CATGCG

CCGTCG-----CGACGG

CCACCC-----GGGTGG

CGATCG-----CGATCG

For use with sequence number 124:

10 CCAAGC-----GCTTGG

CCAAGCC-----GCTTGG

CCAAGCG-----GCGTTGG

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CCAGCG-----CGCTGG

CGATCG-----CGATCG

15 For use with sequence number 125:

CCAAGC------GCTTGG

CGCATG-----CATGCG

CCCAGC------GCTGGG

20 CGATCG-----CGATCG

The use of probe molecules incorporating molecular beacons technology allows for real-time monitoring of amplification reactions, such as NASBA or RT-PCR reactions. The use of molecular beacons technology allows for real-time monitoring of the NASBA reaction (see Leone et al., Nucleic Acids Research., 1998, vol: 26, pp 2150-2155). The molecular beacons probes generally include complementary sequences flanking the HPV-specific sequence, represented herein by the notation arm, and arm, which are capable of hybridising to each other form a stem duplex structure. The precise sequences of arm, and arm, are not material to the invention,

except for the requirement that these sequences must be capable of forming a stem duplex when the probe is not bound to a target HPV sequence.

Molecular beacons probes also include a fluorescent moiety and a quencher moiety, the fluorescent and the quencher moieties being represented herein by the notation X_2 and X_3 . As will be appreciated be the skilled reader, the fluorescer

and quencher moieties are selected such that the quencher moiety is capable of substantially or completely quenching the fluorescence from the fluorescent moiety when the two moieties are in close proximity, e.g. when the probe is in the hairpin

"closed" conformation in the absence of the target sequence. Upon binding to the target sequence, the fluorescent and quencher moieties are held apart such that the fluorescence of the fluorescent moiety is no longer quenched.

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Many examples of suitable pairs of quencher/fluorescer moieties which may be used in accordance with the invention are known in the art (see WO 95/13399, Tyagi and Kramer, ibid). A broad range of fluorophores in many different colours made be used, including for example 5-(2'-aminoethyl)aminonaphthalene-1-sulphonic acid (EDANS), fluorescein, FAM and Texas Red (see Tyagi, Bratu and Kramer, 1998, Nature Biotechnology, 16, 49-53. The use of probes labelled with different coloured fluorophores enables "multiplex" detection of two or more different probes in a single reaction vessel. A preferred quencher is 4-(4'-dimethylaminophenylazo)benzoic acid (DABCYL), a

non-fluorescent chromophore, which serves as a 'universal' quencher for a wide range of fluorophores. The fluorescer and quencher moieties may be covalently attached to the probe in either orientation, either with the fluorescer at or near the 5' end and the quencher at or near the 3' end or vice versa. 5 Protocols for the synthesis of molecular beacon probes are known in the art. A detailed protocol for synthesis is provided in a paper entitled "Molecular Beacons: Hybridization Probes for Detection of Nucleic Acids in Homogenous Solutions" by Sanjay Tyagi et al., 10 Department of Molecular Genetics, Public Health Research Institute, 455 First Avenue, New York, NY 10016, USA, which is available online via the PHRI website (at www.phri.nyu.edu or www.molecular-15 beacons.org).

Suitable combinations of the NASBA P1 and NASBA P2 primer oligonucleotide molecules provided by the invention may be used to drive a NASBA amplification reaction. In order to drive a NASBA amplification reaction the primer 1 and primer 2 oligonucleotides must be capable of priming synthesis of a double-stranded DNA from a target region of mRNA. For this to occur the primer 1 and primer 2 oligonucleotides must comprise target-specific sequences which are complementary to regions of the sense and the antisense strand of the target mRNA, respectively.

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In the first phase of the NASBA amplification cycle, the so-called "non-cyclic" phase, the primer 1 oligonucleotide anneals to a complementary sequence in the target mRNA and its 3' end is extended by the

action of an RNA-dependent DNA polymerase (e.g. reverse transcriptase) to form a first-strand cDNA synthesis. The RNA strand of the resulting RNA: DNA hybrid is then digested, e.g. by the action of RNaseH, to leave a single stranded DNA. The primer 2 oligonucleotide anneals to a complementary sequence towards the 3' end of this single stranded DNA and its 3' end is extended (by the action of reverse transcriptase), forming a double stranded DNA. RNA

polymerase is then able to transcribe multiple RNA copies from the now transcriptionally active promoter sequence within the double-stranded DNA. This RNA transcript, which is antisense to the original target mRNA, can act as a template for a further round of NASBA reactions, with primer 2 annealing to the RNA and priming synthesis of the first cDNA strand and primer 1 priming synthesis of the second cDNA strand. The general principles of the NASBA reaction are well known in the art (see Compton, J. Nature. 350: 91-92).

The target-specific probe oligonucleotides described herein may also be attached to a solid support, such as magnetic microbeads, and used as "capture probes" to immobilise the product of the NASBA amplification reaction (a single stranded RNA). The target-specific "molecular beacons" probes described herein may be used for real-time monitoring of the NASBA reaction.

In a particular embodiment the invention provides the oligonucleotide listed in Table 2, these being NASBA P1 primers and NASBA P2 primers containing the sequences listed in Table 1. The NASBA P1 primers further include a T7 promoter sequence, the NASBA P2

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primers include a sequence for binding of a generic detection probe (see below) and associated probe molecules for use in the detection of HPV mRNA by NASBA. The oligonucleotides listed in Table 2 are merely illustrative and it is not intended that the scope of the invention should be limited to these specific molecules.

The NASBA P2 primers (p2) in Table 2 include the

sequence GATGCAAGGTCGCATATGAG at the 5' end; the NASBA P1

primers (p1) in Table 2 include the sequence

AATTCTAATACGACTCACTATAGGGAGAAGG at the 5' end.

Oligonucleotides suitable for use as probes are

identified by "po". The P2 primers generally contain

HPV sequences from the postive strand, whereas the p1

primers generally contain HPV sequences from the

negative strand. nt-refers to nucleotide position in

the relevant HPV genomic sequence.

20 Table 2-NASBA primer and probe sequences

		HPV	nt
Primer name	Sequence	Type	
I I I I I	TO TO COCCA CAGGAGCGACCC	16	116
HAe6701p2	GATGCAAGGTCGCATATGAGCCACAGGAGCGACCC		
	AGAAAGTTA AATTCTAATACGACTCACTATAGGGAGAAGGACGG	16	368
HAe6701p1	AATTCTAATACGACTCACTATTC		
	TTTGTTGTATTGCTGTTC GATGCAAGGTCGCATATGAGCCACAGGAGCGACCC	16	116
HAe6702p2	GATGCAAGGTCGCATATGAGGGTT		
	AGAAA AATTCTAATACGACTCACTATAGGGAGAAGGGGTT	16	368
HAe6702p1			
	TGTTGTATTGCTGTTC AATTCTAATACGACTCACTATAGGGAGAAGGTCA	16	208
HAe6702Ap1	AATTCTAATACGACTCACTTCACCAGAAGGTTG	16	191
	CGTCGCAGTAACTGT AATTCTAATACGACTCACTATAGGGAGAAGGTTG	1 70	1
HAe6702Bpl		16	186
HAe6702Cp1	AATTCTAATACGACTCACTATAGGGAGANCGT		
111111111111111111111111111111111111111	AGTACACACTTCTA AGTTCTAATACGACTCACTATAGGGAGAAGGGCA	16	185
HAe6702Dpl			114
67007-2	COMCCCATATGAGACAGITAT	. 10	
H16e6702Ap2			
_	GCT		

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	Primer name	Sequence	HPV	nt
		•	Туре	
	H16e6702Bp2	GATGCAAGGTCGCATATGAGATATTAGAATGTGTG	16	182
	H16e6702Cp2	TAC GATGCAAGGTCGCATATGAGTTAGAATGTGTGTAC	16	185
		TGC GATGCAAGGTCGCATATGAGGAATGTGTGTACTGC	16	188
	H16e6702Dp2	AAG		1.40
	H16e6702Apo	ACAGTTATGCACAGAGCT	16	142
5	H16e6702Bpo	ATATTAGAATGTGTGTAC	16	182
•	H16e6702Cpo	TTAGAATGTGTGTACTGC	16	185
	H16e6702Dpo	GAATGTGTGTACTGCAAG	16	188
	HAe67-0-1-po	-GTTTGGTTTTEGGGATTTATGE	-1:6	235
	HAe 6702po	TATGACTTTGCTTTTCGGGA	16	230
10	HAe6702mb1	X ₂ -cgcatgTATGACTTTGCTTTTCGGGAcatgcg	16	230
	HAe6702mb2	$-X_3$ X_2 -ccagctTATGACTTTGCTTTTCGGGAagctgg $-X_3$	16	230
	HAe6702mb3	X ₂ -cacgcTATGACTTTGCTTTTCGGGAgcgtg-X ₃	16	230
15	H16e6702mb4	X ₂ -cgatcgTATGACTTTGCTTTTCGGGAcgatcg -X ₃	16	230
	HAe6703p2	GATGCAAGGTCGCATATGAGCAGAGGAGGAGGATG AAATAGTA	16	656
	HAe6703p1	AATTCTAATACGACTCACTATAGGGAGAAGGGCAC AACCGAAGCGTAGAGTCACAC	16	741
	*** 6702 ·	TGGACAAGCAGAACCGGACAGAGC	16	687
	HAe 6703po HAe 6704p2	GATGCAAGGTCGCATATGAGCAGAGGAGGAGGATG	16	656
20	HAe6704p1	AAATAGA AATTCTAATACGACTCACTATAGGGAGAAGGGCAC	16	741
		AACCGAAGCGTAGAGTCA	16	693
	HAe6704po H18e6701p2	AGCAGAACCGGACAGAGCCCATTA GATGCAAGGTCGCATATGAGACGATGAAATAGATG		702
	H18e6701p1	GAGTT AATTCTAATACGACTCACTATAGGGAGAAGGCACG	18	869
		GACACAAAGGACAG	18	748
25	H18e6701po	AGCCGAACCACAACGTCACA GATGCAAGGTCGCATATGAGGAAAACGATGAAATA		698
	H18e6702p1	GATGGAG AATTCTAATACGACTCACTATAGGGAGAAGGACAC	18	869
		CACGGACACAAAGGACAG		
	H18e6702po	GAACCACAACGTCACACAATG	18	752
	H18e6702pb1	X ₂ -cgcatgGAACCACAACGTCACAATGcatgcg	18	752
	H18e6702mb2	-X ₃ X ₂ -ccgtcgGAACCACAACGTCACACAATGcgacgg	18	. 752
30	H18e6702mb3	-X ₃ . X ₂ -cggaccGAACCACAACGTCACAATGggtccc	j 18	752
	H18e6702mb4	-X ₃ X ₂ -cgatcgGAACCACAACGTCACAATGcgatcg	j 18	752

			HPV	nt
[Primer name	Sequence	Type	
		-X ₃		651
	H18e6703p2	-X ₃ GATGCAAGGTCGCATATGAGTTCCGGTTGACCTTC	18	031
	H1866/0355			817
	H18e6703p1	TATGT AATTCTAATACGACTCACTATAGGGAGAAGGGGTC	18	101/
•	HISEO/OSPI		1	179
	H18e6704p2	GTCTGCTGAGCTTTCT GATGCAAGGTCGCATATGAGGCAAGACATAGAAAT	18	1,3
	HIGEOLOADS		10	379
	H18e6704p1	AATTCTAATACGACTCACTATAGGGAGAAGGACCC	18	
	MIS6010157	AGTGTTAGTTAGTT	1	207
	H18e6704po	TOOR TORONG TO THE CONTROL OF THE CON	31	164
	H31e6701p2	GATGCAAGACAGTATTGGTTGGAGGGAAATACCCTACGA	21	
	noico.orp-			423
	H31e6701p1	TGAAC AATTCTAATACGACTCACTATAGGGAGAAGGGGAC	31	
	Hoteover	ACAACGGTCTTTGACA	31	268
	H31e6701po	T TO COOR CACACACACACACACACACACACACACACACACACACA		164
)	H31e6702p2	GATGCAAGGTCGCATATGAGGGAAATACCCTACGA	.] 31	
,	1.525	TGAACTA	31	423
	H31e6702p1	TGAACTA AATTCTAATACGACTCACTATAGGGAGAAGGCTGG	, 31	
	1132331-1-1	ACACAACGGTCTTTGACA	31	269
	H31e6702po	CCCACACACACACACACACACACACACACACACACA		617
	H31e6703p2	GATGCAAGGTCGCATATGAGACTGACCTCCACTGT	' "	l l
		TATGA	C 31	766
	H31e6703p1	TATGA AATTCTAATACGACTCACTATAGGGAGAAGGTAT	J J*	
		TACTTGTGTGCTCTGT	31	687
.5	H31e6703po	GACAAGCAGAACCGGACACATC		619
	H31e6704p2	GACAAGCAGAACCGGACATATTAGAGTGACCTCCACTGTT GATGCAAGGTCGCATATGAGTGACCTCCACTGTT		
		TGAGCAATT	G 31	766
	H31e6704p1	TGAGCAATT AATTCTAATACGACTCACTATAGGGAGAAGGTGC		1
		AATATCTACTTGTGTGCTCT GT	31	686
	H31e6704po	GGACAAGCAGAACCGGACACATCCAA	31	686
	H31e6704mb	GGACAAGCAGAACCGGACACATCC 1 X ₂ -ccgaaggGACAAGCAGAACCGGACACATCC		
		AAccttcgg -X3	31	686
50	H31e6704mb	AACCTTCGG - X3 2 X2-CCGTCGGGACAAGCAGAACCGGACACATCCA		İ
		Acgacgg -X ₃	31	68
	H31e6704mb	X2- cacgtcgGGACAAGCAGAACCGGACACATCCAA	1	
		•		
		cgacgtg -X ₃ 24 X ₂ -cgcagcGGACAAGCAGAACCGGACACATCCA	A 31	68
	H31e6704mk	1	l l	
		gctgcg-X ₃ o5 X ₂ -cgatcgGGACAAGCAGAACCGGACACATCCA	AA 31	68
	H31e6704m	•	-	
		cgatcg -X₃ 2 GATGCAAGGTCGCATATGAGACTGACCTCCACT	GT 31	61
	. H31e6705p	2 GATGCAAGGTCGCATATGACTOT		
		TAT AATTCTAATACGACTCACTATAGGGAGAAGGCA	ACG 31	80
55	H31e6705p			
		ATTCCAAATGAGCCCAT 2 GATGCAAGGTCGCATATGAGTATCCTGAACCA	ACT 33	63
	H33e6701p			
		GACCTAT		

	Primer name	Sequence	HPV	nt
		_	Type	
	H33e6701p1	AATTCTAATACGACTCACTATAGGGAGAAGGTTGA	33	763
	-	CACATAAACGAACTG		
	H33e6701po	CAGATGGACAAGCACAACC	33	694
	H33e6703p2	GATGCAAGGTCGCATATGAGTCCTGAACCAACTGA	33 .	620
		CCTAT		
	H33e6703p1	AATTCTAATACGACTCACTATAGGGAGAAGGCCCA	33	807
•	•	TAAGTAGTTGCTGTAT		
5	H33e6703po	GGACAAGCACCAGCCACAGC	33	699
-	H33e6703mb1	X2-ccaagcGGACAAGCACAACCAGCCACAGCgct	33	699
	772-6703-62	tgg -X ₃ X ₂ -ccaagcgGACAAGCACAACCAGCCACAGC	33	699
	H33e6703mb2			
		cgcttgg -X ₃ X ₂ -cccagcGGACAAGCACCAGCCACAGCgct	33	699
	H33e6703mb3	X ₂ -cccagcGGACAAGCACAACCAGCCACAGCGCC	33	
		ggg -X ₃		
	H33e6703mb4	X2-ccaaagcGGACAAGCACAACCAGCCACAGCg	33	699
		ctttgg-X ₃		
10	H33e6703mb5	X ₂ -cctgcGGACAAGCACCAGCCACAGCgcagg	33	699
10	H33e0703mb3			
	H33e6703mb6	-X ₃ X ₂ -cgatcgGGACAAGCACAACCAGCCACAGCcga	33	699
	n3366103mm0	N2-CGarcy Ganoral Control of Cont		
		tcg -X ₃	-	421
	H33e6702p2	GATGCAAGGTCGCATATGAGGACCTTTGTGTCCTC	33	431
		AAGAA		618
	H33e6702p1	AATTCTAATACGACTCACTATAGGGAGAAGGAGGT	33	010
۲.		CAGTTGGTTCAGGATA		- 543
-	H33e6702po	AGAAACTGCACTGTGACGTGT	33	543 217
15	H35e6701p2	GATGCAAGGTCGCATATGAGATTACAGCGGAGTGA	35	21/
		GGTAT	25	442
	H35e6701p1	AATTCTAATACGACTCACTATAGGGAGAAGGGTCT	35	442
		TTGCTTTTCAACTGGA	-	070
	H35e5601po	ATAGAGAAGGCCAGCCATAT	35	270 655
	H35e6702p2	GATGCAAGGTCGCATATGAGTCAGAGGAGGAGGAA	35	633
		GATACTA	\	
	H35e6702p1	AATTCTAATACGACTCACTATAGGGAGAAGGGATT	35	844
		ATGCTCTGTGAACA	1	- 610
20	H35e6703p2	GATGCAAGGTCGCATATGAGCCCGAGGCAACTGAC	35	610
		CTATA		
	H35e6703p1	AATTCTAATACGACTCACTATAGGGAGAAGGGTCA	35	770
		ATGTGTGTGCTCTGTA		
	H35e6702po	GACAAGCAAAACCAGACACCTCCAA	35	692
	H35e6703po	GACAAGCAAAACCAGACACC	35	692
	H52e6701p2	GATGCAAGGTCGCATATGAGTTGTGTGAGGTGCTG	52 _	144
		GAAGAAT	<u> </u>	
25	H52e6701p1	AATTCTAATACGACTCACTATAGGGAGAAGGCCCT	52	358
		CTCTTCTAATGTTT		
	H52e6701po	GTGCCTACGCTTTTTATCTA	52	296
	H52e6702p2	GATGCAAGGTCGCATATGAGGTGCCTACGCTTTTT	52	296

			HP	J	nt
Γ	Primer name	Sequence	Тур	pe]	
1					
F		ATCTA	52		507
ŀ	H52e6702p1	ATCTA AATTCTAATACGACTCACTATAGGGAGAAGGGGGG		1	
1		TCTCCAACACTCTGAACA	52		461
- }	H52e6702po		58		157
-	H58e6701p2	TGCAAACAAGCGATTICA GATGCAAGGTCGCATATGAGTCAGGCGTTGGAGAC	"	1	
1	HOSEGIOTPE		58	,	301
	0001 1	ATC AATTCTAATACGACTCACTATAGGGAGAAGGAGCA	20)	302
	H58e6701p1	T T C T C T			173
		ATCGTAAGCACACT GATGCAAGGTCGCATATGAGTCTGTGCATGAAATC	58	3	1/3
	H58e6702p2	GATGCAAGGICGC	1		
		GAA AATTCTAATACGACTCACTATAGGGAGAAGGAGCA	5	8	291
	H58e6702p1				1
		CACTTTACATACTG	15	8	192
	H58e6701po	TCAAATGCGTTGAATGCA		8	218
	H58e6702po		1 -	(11)	514
	HBe6701p2	TTGCAGCGATCTGAGGTATATCACTGCTGGACAA	, _	,,,	
	nbed/01P1	ì		3(11)	619
	0703.1	CAT AATTCTAATACGACTCACTATAGGGAGAAGGTCAT	- -	3(11)	1 3
	HBe6701p1		- 1		+ -14
		CTTCTGAGCTGTCT GATGCAAGGTCGCATATGAGTACACTGCTGGACAA	$A \mid I$	B(11)	514
	HBe6702p2				
		CATGCA AATTCTAATACGACTCACTATAGGGAGAAGGGTC	A	B(11)	693
	HBe6702p1	AATTCTAATACGACTCACTATAGGGACT	1		
		CATCCACAGCAACAGGTCA		B(11)	59
	HBe6701po	GTAGGGTTACATTGCTATGA		B(11)	59
	НВе6702ро			B(11)	69
	HBe6703p2	GTAGGGTTACATTGCTATGAGGGTGACCTGTTGCTGT GATGCAAGGTCGCATATGAGTGACCTGTTGCTGT	9	- (\
	nbec705P2		1	B(11)	83
	6703-1	GATGTGA AATTCTAATACGACTCACTATAGGGAGAAGGTAC		D(11)	
ı	HBe6703pl	TGAATCGTCCGCCAT			79
				B(11)	29
	HBe6703po	ATWGTGTGTCCCATCIGC GATGCAAGGTCGCATATGAGCATGCCATAAATG	TA	C(18	l
5	HCe6701p2	GATGCAAGGICGC		39 45)	
		TAGA AATTCTAATACGACTCACTATAGGGAGAAGGCA	CC	C(18	40
	HCe6701p1	AATTCTAATACGACTCACTATAGG		39 45	
		GCAGGCACCTTATTAA		C(18	3
	HCe6701po	-	1	39 45	
	11000111		17) C	39	2
	H39e6701p	2 GATGCAAGGTCGCATATGAGGCAGACGACCACT	AC		.
	H3960,015			1-20	- 1 3
		TRACAR CONCERN TO A COLOR CONCERN TO A COLOR COL	CAC	39	
	H39e6701p				
		CGAGTCCGAGTAATA		39	2
0	H39e6701p	ATAGGACGGGAACCACT ATAGGACGGCGCATATGAGTATTACTCGGAC ATAGGAAGGTCGCATATGAGTATTACTCGGAC	TCG	39]
-	H39e6702p	2 GATGCAAGGTCGCATATGAGTTT			
			TTG	39	
	H39e6702p	AATTCTAATACGACTCACTATAGGGAGAAGG			
	1139607521	CGTTTCTCTTCGTGTTA		39	
	1120 6700		CCC		
	H39e6702r	TOTAL TOTAL CONCCCATA TOTAL CONCENTRATION OF THE PROPERTY OF T		' "	1
	H39e6703	PZ GAIGOIZII			

	Primer name	Sequence	HPV	nt
			Туре	200
	H39e6703p1	AATTCTAATACGACTCACTATAGGGAGAAGGGCAC	39	886
		ACCACGGACACAAA		
	H39e6703po	TAGCCAGACGGGATGAACCACAGC	39	749
	H45e6701p2	GATGCAAGGTCGCATATGAGAACCATTGAACCCAG	45	430
		CAGAAA		
	H45e6701p1	AATTCTAATACGACTCACTATAGGGAGAAGGTCTT	45	527
		TCTTGCCGTGCCTGGTCA		
5	H45e6702p2	GATGCAAGGTCGCATATGAGGAAACCATTGAACCC	45	428
		AGCAGAAAA		
	H45e6702p1	AATTCTAATACGACTCACTATAGGGAGAAGGTTGC	45	558
		TATACTTGTGTTTCCCTACG		
	H45e6701po	GTACCGAGGGCAGTGTAATA	45	500
	H45e6702po	GGACAAACGAAGATTTCACA	45	467
	H45e6703p2	GATGCAAGGTCGCATATGAGGTTGACCTGTTGTGT	45	656
	114360,002	TACCAGCAAT		
10	H45e6703pl	AATTCTAATACGACTCACTATAGGGAGAAGGCACC	45	8 68
10	n43e0703pi	ACGGACACAAAGGACAAG		
	H45e6704p2	GATGCAAGGTCGCATATGAGCTGTTGACCTGTTGT	45	654
	H45e6/U4p2			
		GTTACGA AATTCTAATACGACTCACTATAGGGAGAAGGCCAC	45	868
	H45e6704pl			Ì
		GGACACAAAGGACAAG GATGCAAGGTCGCATATGAGGTTGACCTGTTGTGT	45	656
	H45e6705p2	GATGCAAGGTCGCATATGAGGTTGACCTGTTGTGT		
		TACGA	45	868
	H45e6705p1	AATTCTAATACGACTCACTATAGGGAGAAGGACGG	13	"
		ACACACAAAGGACAAG	45	686
15	H45e6703po	GAGTCAGAGGAGGAAAACGATG	45	696
	H45e6704po	AGGAAAACGATGAAGCAGATGGAGT	45	730
	H45e6705po	ACAACTACCAGCCGACGAGCCGAA	51	658
	H51e6701p2	GATGCAAGGTCGCATATGAGGGAGGAGGATGAAGT	31	
		AGATA	51	807
	H51e6701p1	AATTCTAATACGACTCACTATAGGGAGAAGGGCCC	31	00,
		ATTAACATCTGCTGTA	 	655
20	H51e6702p2	GATGCAAGGTCGCATATGAGAGAGGAGGAGGATGA	51	655
		AGTAGATA		
	H51e6702p1	AATTCTAATACGACTCACTATAGGGAGAAGGACGG	51	829
		GCAAACCAGGCTTAGT		
	H51e6701po	GCAGGTGTTCAAGTGTAGTA	51	74
	H51e6702po	TGGCAGTGGAAAGCAGTGGAGACA	51	77:
	H56e6701p2	GATGCAAGGTCGCATATGAGTTGGGGTGCTGGAGA	56	519
		CAAACATCT		
25	H56e6701p1	AATTCTAATACGACTCACTATAGGGAGAAGGTTCA	56	66
20		TCCTCATCCTCATCCTCTGA		
	H56e6702p2	GATGCAAGGTCGCATATGAGTGGGGTGCTGGAGAC	56	52
	11306070252	AAACATC	L	
	TIE C - 6702-1	AATTCTAATACGACTCACTATAGGGAGAAGGCATC	56	66
	H56e6702p1	CTCATCCTCATCCTCTGA		
	H56e6703p2		56	51

			HPV	nt
	Primer name	Sequence	Type	
		ON A TOTAL		
		CAAACAT AATTCTAATACGACTCACTATAGGGAGAAGGCCAC	56	764
	H56e6703p1			
		AAACTTACACTCACAACA AAAGTACCAACGCTGCAAGACGT	56	581
30	H56e6701po	AGAACTACCAACGCTGCAAACAGAAAT AGAACTAACACCTCAAACAGAAAT	56	610
	H56e6702po	AGAACTAACACCTCAAACACTTT AGTACCAACGCTGCAAGACGTT	56	583
	H56e6703po	TTGGACAGCTCAGAGGATGAGG	56	656
	H56e6703po1	GATGCAAGGTCGCATATGAGGATTTTCCTTATGCA	56	279
	H56e6704p2	- 1		
		GTGTG AATTCTAATACGACTCACTATAGGGAGAAGGGACA	56	410
35	H56e6704pl	L Company of the Comp		
		TCTGTAGCACCTTATT	56	348
	H56e6704po	GACTATTCAGTGTATGGAGC	A (16	
	HPVAPO1A	CAACTGAYCTMYACTGTTATGA	31 35)	
			A (16	1
	HPVApo1Amb1	X2-cgcatgCAACTGAYCTMYACTGTTATGAcatgcg	1	1
		1. 17	31 35)	1
	HPVApolAmb2	X_2 -ccgtcgCAACTGAYCTMYACTGTTATGAcga	A (16	
	III VIADOLI ZIII		31 35)	
		cgg -X₃ X₂-ccacccCAACTGAYCTMYACTGTTATGAgg	A (16	
40	HPVApo1Amb3	X2-ccacccCAACTGAICIMIACIGIMITOTIS	31 35)	
		gtgg -X ₃	1	
	HPVApolAmb4	gtgg - X ₃ X ₂ -cgatcgCAACTGAYCTMYACTGTTATGAcga	A (16	1 1
	NE AMPOITMENT		31 35)	
		tcg-X ₃	A (33	
	HPVAPO4A	GAAMCAACTGACCTAYWCTGCTAT	52 58)	1
		TOTAL COMPANIE CONTRACTOR AND CONTRA	A (33	1
	HPVAPO4Amb1	X2-ccaagcGAAMCAACTGACCTAYWCTGCTATgc	52 58)	
		ttgg-X ₃	_ \	
	HPVAPO4Amb2		A (33	
	HPVAPOTAMOZ		52 58)	
		ggcttgg -X3	A (33	-
45	HPVAPO4Amb3	ggcccgg X3 X2-ccaagcgGAAMCAACTGACCTAYWCTGCTA	52 58)	
		Tcgcttgg -X ₃	1	
	HPVAPO4Amb4		A (33	}
	HEVALOGAMOS		52 58)	1
		ctgg -X ₃	A (33	
	HPVAPO4Amb	Ctgg -x ₃ X ₂ -cgatcgGAAMCAACTGACCTAYWCTGCTATcg	52 58)	}
		atcg-X ₃		
	HPVCPO4	AAGACATTATTCAGACTC	C (18	
	HPVCF04	12.01.01.0	45 39)	
		1 X ₂ -ccaagcAAGACATTATTCAGACTCgcttgg-X	C (18	
	HPVCPO4Amb	X ₂ -ccaagcaagcaacaiiiiiio	45 39)	
		2 X ₂ -cgcatgAAGACATTATTCAGACTCcatgcg-X	C (18	
50	HPVCPO4Amb	Z X2-cgcatgAAGACATTAIICAGACICCatgcg	45 39)	
		The state of the s		
	HPVCP04Amb	3 X ₂ -cccagcAAGACATTATTCAGACTCgctggg->	45 39)	
				_
	HPVCPO4Amb	4 X2-cgatcgAAGACATTATTCAGACTCcgatcg-X	45 39)	
			45 35)	

The meaning of X_2- and $-X_3$ is defined above, in the discussion of "molecular beacons" probe molecules.

In a further embodiment the invention provides
the oligonucleotides listed in Table 3, these being
PCR primers for use in the detection of HPV mRNA by
RT-PCR. These oligonucleotides are merely
illustrative and it is not intended that the scope of
the invention should be limited to these specific

molecules:

	Primer name	Sequence	HPV type	nt
	77 . C701 DCD2	CCACAGGAGCGACCCAGAAAGTTA	16	116
	HAe6701PCR2	ACGGTTTGTTGTATTGCTGTTC	16	368
4 -	HAe 6701 PCR1	CCACAGGAGCGACCCAGAAA	16	116
15	HAe6702PCR2	GGTTTGTTGTATTGCTGTTC	16	368
	HAe6702PCR1	CAGAGGAGGAGGATGAAATAGTA	16	656
	HAe6703PCR2	GCACAACCGAAGCGTAGAGTCACAC	16	741
	HAe 6703 PCR1	CAGAGGAGGAGGATGAAATAGA	16	656
	HAe6704PCR2	GCACAACCGAAGCGTAGAGTCA	16	741
20	HAe6704PCR1	ACGATGAAATAGATGGAGTT	18	702
,	H18e6701PCR2	CACGGACACACAAAGGACAG	18	869
	H18e6701PCR1	GAAAACGATGAAATAGATGGAG	18	698
	H18e6702PCR2	ACACCACGGACACACAAAGGACAG	18	869
	H18e6702PCR1	TTCCGGTTGACCTTCTATGT	18	651
25	H18e6703PCR2	GGTCGTCTGCTGAGCTTTCT	18	817
	H18e6703PCR1	GCAAGACATAGAAATAACCTG	18	179
	H18e6704PCR2		18	379
	H18e6704PCR1	ACCCAGTGTTAGTTAGTT GGAAATACCCTACGATGAAC	31	164
	H31e6701PCR2	GGACACACGGTCTTTGACA	31	423
30	H31e6701PCR1		31	164
	H31e6702PCR2	GGAAATACCCTACGATGAACTA	31	423
	H31e6702PCR1	CTGGACACAACGGTCTTTGACA	31	617
	H31e6703PCR2	ACTGACCTCCACTGTTATGA	31	766
	H31e6703PCR1	TATCTACTTGTGTGCTCTGT	31	619
35	H31e6704PCR2	TGACCTCCACTGTTATGAGCAATT	31	.766
	H31e6704PCR1	TGCGAATATCTACTTGTGTGCTCT GT	31	617
	H31e6705PCR2	ACTGACCTCCACTGTTAT	31	809
	H31e6705PCR1	CACGATTCCAAATGAGCCCAT	33	618
	H33e6701PCR2	TATCCTGAACCAACTGACCTAT	33	763
40	H33e6701PCR1	TTGACACATAAACGAACTG	33	620
	H33e6703PCR2	TCCTGAACCAACTGACCTAT	33	807
	H33e6703PCR1	CCCATAAGTAGTTGCTGTAT		431
	H33e6702PCR2	GACCTTTGTGTCCTCAAGAA	33	618
	H33e6702PCR1	AGGTCAGTTGGTTCAGGATA	33	217
45	H35e6701PCR2	ATTACAGCGGAGTGAGGTAT	35	
	H35e6701PCR1	GTCTTTGCTTTTCAACTGGA	35	442
	H35e6702PCR2	TCAGAGGAGGAAGATACTA	35	655

		2 200	HPV	nt
Γ	Primer name	Sequence	type	844
		GATTATGCTCTCTGTGAACA	35	
Γ	H35e6702PCR1	CCCGAGGCAACTGACCTATA	35	610
Ī	H35e6703PCR2	GTCAATGTGTGTGCTCTGTA	35	770
Ī	H35e6703PCR1	TTGTGAGGTGCTGGAAGAAT	52	144
Ī	H52e6701PCR2	CCCTCTCTAATGTTT	52	358
5	H52e6701PCR1	GTGCCTACGCTTTTTATCTA	52	296
	H52e6702PCR2	GGGGTCTCCAACACTCTGAACA	52	507
	H52e6702PCR1	GGGTCTCCAACACTGTG	58	157
	H58e6701PCR2	TCAGGCGTTGGAGACATC	58	301
	H58e6701PCR1	AGCAATCGTAAGCACACT	58	173
0	H58e6702PCR2	TCTGTGCATGAAATCGAA	58	291
J	H58e6702PCR1	AGCACACTTTACATACTG	B(11)	514
	HBe6701PCR2	TACACTGCTGGACAACAT	B(11)	619
	HBe6701PCR1	TCATCTTCTGAGCTGTCT	B(11)	514
	HBe6702PCR2	TACACTGCTGGACAACATGCA	B(11)	693
_	HBe6702PCR1	GTCACATCCACAGCAACAGGTCA	B(11)	693
5	HBe6703PCR2	TGACCTGTTGCTGTGGATGTGA	B(11)	832
	HBe6703PCR1	TACCTGAATCGTCCGCCAT	C (18	295
	HCe6701PCR2	CATGCCATAAATGTATAGA	39 45	
	HC66/01FCKZ		C (18	408
	HCe6701PCR1	CACCGCAGGCACCTTATTAA	39 45	
	HC6010TLCKT		39	210
	H39e6701PCR2	GCAGACGACCACTACAGCAAA	39	344
20	H39e6701PCR1	ACACCGAGTCCGAGTAATA	39	344
	H39e6701PCR2	TATTACTCGGACTCGGTGT	39	558
	H39e6702PCR1	CTTGGGTTTCTCTTCGTGTTA	39	703
	H39e6702FCR1	CAAATAGATGAACCCGACCA	39	886
	H39e6703PCR2	CCACACCACGGACACACAAA	45	430
25	H39e6703PCR1	AACCATTGAACCCAGCAGAAA	45	527
	H45e6701PCR2	TOTTTCTTGCCGTGCCTGGTCA	45	428
	H45e6701PCR1	CADACCATTGAACCCAGCAGAAAA	45	558
	H45e6702PCR2	TTCCTATACTTGTGTTTCCCTACG		656
	H45e6702PCR1	CUTCACCTGTTGTGTTACCAGCAAT	45	868
30	H45e6703PCR2	CACCACGGACACACAAAGGACAAG	45	654
	H45e6703PCR1	CTGTTGACCTGTTGTGTTACGA	45	868
	H45e6704PCR2	CCACGGACACACAAAGGACAAG	45	656
	H45e6704PCR1	GTTGACCTGTTGTGTTACGA	45	
	H45e6705PCR2	ACGGACACAAAGGACAAG	45	
35	H45e6705PCR1	GGAGGAGGATGAAGTAGATA	51	658
	H51e6701PCR2	GCCCATTAACATCTGCTGTA	51	807
	H51e6701PCR1	AGAGGAGGAGGATGAAGTAGATA	51	655
	H51e6702PCR2	TO CONTRACTOR	51	829
	H51e6702PCR1	- CACACATCI	56	519
40	H56e6701PCR2	TTGGGGTGCTGGAGACTCTCTGA	56	665
	H56e6701PCR1	TTCATCCTCATCCTCAAACATC	56	520
	H56e6702PCR2	TOCCCCT GCTGGAGACTATION	56	665
	H56e6702PCR1	CATCUICATCCTCATCA	56	519
	H56e6703PCR2	T TOTAL TOTA	56	764
1 ⊑	H56e6703PCR1	CCACAAACTTACACTCACTC	56	279
45	H56e6704PCR2	CATHUTCCTTATGCAGTOT	56	410
	H56e6704PCR			

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Primer-pairs and primer-probe sets

The invention further provides primer-pairs and primer/probe sets for use in the detection of HPV E6 transcripts.

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A "primer-pair" is taken to mean two primers which may be used in combination for amplification of a portion of an HPV E6 transcript, for example by NASBA or RT-PCR. The individual oligonucleotide primers making up the primer-pair may be supplied separately, e.g. in separate containers. A primer-pair may also be supplied as a homogenous mixture of the two primers, this mixture may include additional reagents required for the amplification reaction, as discussed below.

A "primer/probe set" is taken to mean a set of oligonucleotides comprising a primer-pair, as defined above, and at least one oligonucleotide probe which is suitable for use in detection of an amplification product generated by use of the primer-pair. The individual oligonucleotides making up the primer/probe set may be supplied separately, e.g. in separate containers or as a homogenous mixture.

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In this context "primer" is taken to encompass primers suitable for use in PCR and primers suitable for use in NASBA.

The term "probe" may encompass any of the probe types described herein, including molecular beacons probes suitable for use in real-time NASBA (see below) and capture probes for immobilisation of NASBA amplification products.

Specific primer-pairs provided by the invention are given below, together with suitable probes which may be used in the detection of amplification products

generated using the primer-pair. In preferred embodiments, the primer-pairs listed below may comprise a NASBA P1 primer and a NASBA P2 primer or two PCR primers. The most preferred specific primer combinations are listed, using the primer names given in Tables 2 and 3. However, it is not intended to limit the scope of the invention to these particular combinations:

10 Primer-pairs and probes for use in the detection of mRNA transcripts from the E6 gene of HPV 16:

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(1) an oligonucleotide primer comprising sequence number 1 and an oligonucleotide primer comprising sequence number 2; oligonucleotide probe comprising sequence number 5.

Preferred NASBA primers: HAe6701p1 and HAe6701p2 Preferred PCR primers: HAe6701PCR1 and HAe6701PCR2

(2) an oligonucleotide primer comprising sequence number 3 and an oligonucleotide primer comprising sequence number 4; oligonucleotide probe comprising sequence number 6.

Preferred NASBA primers: HAe6702p1 and HAe6702p2 Preferred PCR primers: HAe 6702PCR1 and HAe6702PCR2

(3) an oligonucleotide primer comprising sequence number 7 and an oligonucleotide primer comprising sequence number 8; oligonucleotide probe comprising sequence number 9.

Preferred NASBA primers: HAe6703pl and HAe6703p2

Preferred PCR primers: HAe6703PCR1 and HAe6703PCR2

(4) an oligonucleotide primer comprising sequence number 10 and an oligonucleotide primer comprising sequence number 11; oligonucleotide probe comprising sequence number 12.

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Preferred NASBA primers: HAe6704p1 and HAe6704p2 Preferred PCR primers: HAe6704PCR1 and HAe6704PCR2

- (5) an oligonucleotide primer comprising one of

 10 sequence numbers 126, 127, 128 or 129 and an
 oligonucleotide primer comprising sequence number 1 or
 sequence number 3.
- (6) an oligonucleotide primer comprising sequence number 2 or sequence number 4 and an oligonucleotide primer comprising one of sequence numbers 130, 131, 132 or 133.
- Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 18:
- (7) an oligonucleotide primer comprising sequence number 13 and an oligonucleotide primer comprising sequence number 14; oligonucleotide probe comprising sequence number 15.

Preferred NASBA primers: H18e6701p1 and H18e6701p2 Preferred PCR primers: H18e6701PCR1 and H18e6701PCR2

- 30 (8) an oligonucleotide primer comprising sequence number 16 and an oligonucleotide primer comprising sequence number 17; oligonucleotide probe comprising sequence number 18.
- Preferred NASBA primers: H18e6702p1 and H18e6702p2 Preferred PCR primers: H18e6702PCR1 and H18e6702PCR2

- (9) an oligonucleotide primer comprising sequence number 19 and an oligonucleotide primer comprising sequence number 20.
- Preferred NASBA primers: H18e6703pl and H18e6703p2 Preferred PCR primers: H1836703PCR1 and H18e6703PCR2
- (10) an oligonucleotide primer comprising sequence number 21 and an oligonucleotide primer comprising sequence number 22; oligonucleotide probe comprising sequence number 23.

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Preferred NASBA primers: H18e6704pl and H18e6704p2 Preferred PCR primers: H18e6704PCR1 and H18e6704PCR2

Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 31:

- (11) an oligonucleotide primer comprising sequence number 24 and an oligonucleotide primer comprising sequence number 25; oligonucleotide probe comprising sequence number 26.
- Preferred NASBA primers: H31e6701p1 and H31e6701p2

 Preferred PCR primers: H31e6701PCR1 and H31e6701PCR2
 - (12) an oligonucleotide primer comprising sequence number 27 and an oligonucleotide primer comprising sequence number 28; oligonucleotide probe comprising sequence number 29.

Preferred NASBA primers: H31e6702p1 and H31e6702p2 Preferred PCR primers: H31e6702PCR1 and H3136702PCR2

35 (13) an oligonucleotide primer comprising sequence number 30 and an oligonucleotide primer comprising

sequence number 31; oligonucleotide probe comprising sequence number 32.

- Preferred NASBA primers: H31e6703pl and H31e6703p2

 Preferred PCR primers: H31e6703PCR1 and H31e6703PCR2
 - (14) an oligonucleotide primer comprising sequence number 33 and an oligonucleotide primer comprising sequence number 34; oligonucleotide probe comprising
- 10 sequence number 35.

Preferred NASBA primers: H31e6704p1 and H31e6704p2 Preferred PCR primers: H31e6704PCR1 and H312e6704PCR2

- 15 (15) an oligonucleotide primer comprising sequence number 36 and an oligonucleotide primer comprising sequence number 37;
- Preferred NASBA primers: H31e6705p1 and H31e6705p2

 Preferred PCR primers: H31e6705PCR1 and H31e6705PCR2

Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 33:

- 25 (16) an oligonucleotide primer comprising sequence number 38 and an oligonucleotide primer comprising sequence number 39; oligonucleotide probe comprising sequence number 40.
- Preferred NASBA primers: H33e6701p1 and H33e6701p2 Preferred PCR primers: H33e6701PCR1 and H33e6701PCR2
- (17) an oligonucleotide primer comprising sequence number 41 and an oligonucleotide primer comprising sequence number 42; oligonucleotide probe comprising sequence number 43.

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Preferred NASBA primers: H33e6703pl and H33e6703p2 Preferred PCR primers: H33e6703PCR1 and H33e6703PCR2

- (18) an oligonucleotide primer comprising sequence number 44 and an oligonucleotide primer comprising sequence number 45; oligonucleotide probe comprising sequence number 46.
- Preferred NASBA primers: H33e6702p1 and H33e6702p2

 Preferred PCR primers: H33e6702PCR1 and H33e6702PCR2

Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 35:

- 15 (19) an oligonucleotide primer comprising sequence number 47 and an oligonucleotide primer comprising sequence number 48; oligonucleotide probe comprising sequence number 53.
- Preferred NASBA primers: H35e6701p1 and H35e6701p2
 Preferred PCR primers: H35e6701PCR1 and H35e6701PCR2
- (20) an oligonucleotide primer comprising sequence number 49 and an oligonucleotide primer comprising sequence number 50; oligonucleotide probe comprising sequence number 54.

Preferred NASBA primers: H35e6702p1 and H35e6702p2 Preferred PCR primers: H35e6702PCR1 and H35e6702PCR2

(21) an oligonucleotide primer comprising sequence number 51 and an oligonucleotide primer comprising sequence number 52; oligonucleotide probe comprising sequence number 55.

Preferred NASBA primers: H35e6703pl and H35e6703p2 Preferred PCR primers: H35e6703PCR1 and H35e6703PCR2

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Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 52:

(22) an oligonucleotide primer comprising sequence number 56 and an oligonucleotide primer comprising sequence number 57; oligonucleotide probe comprising sequence number 58.

Preferred NASBA primers: H52e6701p1 and H52e6701p2

10 Preferred PCR primers: H52e6701PCR1 and H52e6701PCR2

(23) an oligonucleotide primer comprising sequence number 59 and an oligonucleotide primer comprising sequence number 60; oligonucleotide probe comprising sequence number 61.

Preferred NASBA primers: H52e6702p1 and H52e6702p2 Preferred PCR primers: H52e6702PCR1 and H52e6702PCR2

- 20 Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 58:
- (24) an oligonucleotide primer comprising sequence number 62 and an oligonucleotide primer comprising
 sequence number 63; oligonucleotide probe comprising sequence number 66.

Preferred NASBA primers: H58e6701p1 and H58e6701p2 Preferred PCR primers: H58e6701PCR1 and H58e6701PCR2

(25) an oligonucleotide primer comprising sequence number 64 and an oligonucleotide primer comprising sequence number 65; oligonucleotide probe comprising sequence number 67.

Preferred NASBA primers: H58e6702p1 and H58e6702p2 Preferred PCR primers: H58e6702PCR1 and H58e6702PCR2

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Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 51:

(26) an oligonucleotide primer comprising sequence number 104 and an oligonucleotide primer comprising sequence number 105; oligonucleotide probe comprising sequence number 108.

Preferred NASBA primers: H51e6701p1 and H51e6701p2

Preferred PCR primers: H51e6701PCR1 and H51e6701PCR2

(27) an oligonucleotide primer comprising sequence number 106 and an oligonucleotide primer comprising sequence number 107; oligonucleotide probe comprising sequence number 109.

Preferred NASBA primers: H51e6702p1 and H51e6702p2 Preferred PCR primers: H51e6702PCR1 and H51e6702PCR2

20 Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 56:

(28) an oligonucleotide primer comprising sequence number 110 and an oligonucleotide primer comprising sequence number 111; oligonucleotide probe comprising sequence number 116.

Preferred NASBA primers: H56e6701p1 and H56e6701p2 Preferred PCR primers: H56e6701PCR1 and H56e6701PCR2

(29) an oligonucleotide primer comprising sequence number 112 and an oligonucleotide primer comprising sequence number 113; oligonucleotide probe comprising sequence number 117.

Preferred NASBA primers: H56e6702p1 and H56e6702p2 Preferred PCR primers: H56e6702PCR1 and H56e6702PCR2

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(30) an oligonucleotide primer comprising sequence number 114 and an oligonucleotide primer comprising sequence number 115; oligonucleotide probe comprising sequence number 118 or sequence number 119.

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Preferred NASBA primers: H56e6703p1 and H56e6703p2 Preferred PCR primers: H56e6703PCR1 and H56e6703PCR2

(31) an oligonucleotide primer comprising sequence

-10 number 120 and an oligonucleotide primer comprising
sequence number 121; oligonucleotide probe comprising
sequence number 122.

Preferred NASBA primers: H56e6704p1 and H56e6704p2

Preferred PCR primers: H56e6704PCR1 and H56e6704PCR2

Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 39:

- 20 (32) an oligonucleotide primer comprising sequence number 80 and an oligonucleotide primer comprising sequence number 81; oligonucleotide probe comprising sequence number 82.
- Preferred NASBA primers: H39e6701p1 and H39e6701p2
 Preferred PCR primers: H39e6701PCR1 and H39e6701PCR2
- (33) an oligonucleotide primer comprising sequence number 83 and an oligonucleotide primer comprising sequence number 84; oligonucleotide probe comprising sequence number 85.

Preferred NASBA primers: H39e6702p1 and H39e6702p2 Preferred PCR primers: H39e6702PCR1 and H39e6702PCR2

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(34) an oligonucleotide primer comprising sequence number 86 and an oligonucleotide primer comprising

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sequence number 87; oligonucleotide probe comprising sequence number 88.

Preferred NASBA primers: H39e6703p1 and H39e6703p2

Preferred PCR primers: H39e6703PCR1 and H39e6703PCR2

Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of HPV 45:

- 10 (35) an oligonucleotide primer comprising sequence number 89 and an oligonucleotide primer comprising sequence number 90; oligonucleotide probe comprising sequence number 93.
- Preferred NASBA primers: H45e6701pl and H45e6701p2
 Preferred PCR primers: H45e6701PCR1 and H45e6701PCR2
- (36) an oligonucleotide primer comprising sequence number 91 and an oligonucleotide primer comprising
 sequence number 92; oligonucleotide probe comprising sequence number 94.

Preferred NASBA primers: H45e6702p1 and H45e6702p2 Preferred PCR primers: H45e6702PCR1 and H45e6702PCR2

(37) an oligonucleotide primer comprising sequence number 95 and an oligonucleotide primer comprising sequence number 96; oligonucleotide probe comprising sequence number 101.

Preferred NASBA primers: H45e6703p1 and H45e6703p2 Preferred PCR primers: H45e6703PCR1 and H45e6703PCR2

(38) an oligonucleotide primer comprising sequence number 97 and an oligonucleotide primer comprising sequence number 98; oligonucleotide probe comprising sequence number 102.

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Preferred NASBA primers: H45e6704p1 and H45e6704p2 Preferred PCR primers: H45e6704PCR1 and H45e6704PCR2

(39) an oligonucleotide primer comprising sequence number 99 and an oligonucleotide primer comprising sequence number 100; oligonucleotide probe comprising sequence number 103.

Preferred NASBA primers: H45e6705p1 and H45e6705p2

OPREFERRED PCR primers: H45e6705PCR1 and H45e6705PCR2

Primer-pairs for use in the detection of mRNA transcripts from the E6 gene of group B HPV:

- 15 (40) an oligonucleotide primer comprising sequence number 68 and an oligonucleotide primer comprising sequence number 69; oligonucleotide probe comprising sequence number 72.
- Preferred NASBA primers: HBe6701p1 and HBe6701p2 Preferred PCR primers: HBe6701PCR1 and HBe6701PCR2
- (41) an oligonucleotide primer comprising sequence number 70 and an oligonucleotide primer comprising
 25 sequence number 71; oligonucleotide probe comprising sequence number 73.

Preferred NASBA primers: HBe6702p1 and HBe6702p2 Preferred PCR primers: HBe6702PCR1 and HBe6702PCR2

(42) an oligonucleotide primer comprising sequence number 74 and an oligonucleotide primer comprising sequence number 75; oligonucleotide probe comprising sequence number 76.

Preferred NASBA primers: HBe6703p1 and HBe6703p2 Preferred PCR primers: HBe6703PCR1 and HBe6703PCR2

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Primer-pair for use in the detection of mRNA transcripts from the E6 gene of group C HPV:

(43) an oligonucleotide primer comprising sequence number 77 and an oligonucleotide primer comprising sequence number 78; oligonucleotide probe comprising sequence number 79.

Preferred NASBA primers: HCe6701p1 and HCe6701p2

Preferred PCR primers: HCe6701PCR1 and HCe6701PCR2

Methods of detecting HPV

In a further aspect the invention provides a method for detecting HPV mRNA in a test sample suspected of containing HPV which comprises performing an amplification reaction on the test sample to amplify a portion of the mRNA transcribed from the E6 gene of HPV, wherein the amplification reaction is performed using one of the primer-pairs provided by the invention, as defined above.

Preferred amplification techniques which may be used to amplify a portion of the E6 mRNA are RT-PCR or NASBA.

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The "test sample suspected of containing HPV" will most commonly be a clinical sample, for example a cervical scraping in the cervical screening field. The amplification reaction will preferably be carried out on a preparation of nucleic acid isolated from the test sample. The preparation of nucleic acid must include mRNA, however it need not be a preparation of purified poly A+ mRNA and preparations of total RNA or crude preparations of total nucleic acid containing both RNA and genomic DNA are also suitable as starting material for a NASBA reaction. Essentially any technique known in the art for the isolation of a

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preparation of nucleic acid including mRNA may be used to isolate nucleic acid from the test sample. A preferred technique is the "Boom" isolation method described in US-A-5,234,809 and EP-B-0389,063. This method, which can be used to isolate a nucleic acid preparation containing both RNA and DNA, is based on the nucleic acid binding properties of silicon dioxide particles in the presence of the chaotropic agent guanidine thiocyanate (GuSCN).

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Methods for the detection of HPV in a test sample using the NASBA technique will generally comprise the following steps:

- (a) assembling a reaction medium comprising a

 primer-pair according to the invention, an RNA
 directed DNA polymerase, a ribonuclease that
 hydrolyses the RNA strand of an RNA-DNA hybrid without
 hydrolysing single or double stranded RNA or DNA, an
 RNA polymerase that recognises said promoter, and
 ribonucleoside and deoxyribonucleoside triphosphates;
 - (b) incubating said reaction medium with a preparation of nucleic acid isolated from a test sample suspected of containing HPV under reaction conditions which permit a NASBA amplification reaction; and
 - (c) detecting and/or quantitatively measuring any HPV-specific product of the NASBA amplification reaction.
- Detection of the specific product(s) of the NASBA reaction (i.e. sense and/or antisense copies of the target RNA) may be carried out in a number of different ways. In one approach the NASBA product(s) may be detected with the use of an HPV-specific hybridisation probe capable of specifically annealing to the NASBA product. The hybridisation probe may be attached to a revealing label, for example a

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fluorescent, luminescent, radioactive or chemiluminescent compound or an enzyme label or any other type of label known to those of ordinary skill in the art. The precise nature of the label is not critical, but it should be capable of producing a signal detectable by external means, either by itself or in conjunction with one or more additional substances (e.g. the substrate for an enzyme).

Also within the scope of the invention is so-10 called "real-time NASBA" which allows continuous monitoring of the formation of the product of the NASBA reaction over the course of the reaction. In a preferred embodiment this may be achieved using a "molecular beacons" probe comprising an HPV-specific 15 sequence capable of annealing to the NASBA product, a stem-duplex forming oligonucleotide sequence and a pair of fluorescer/quencher moieties, as known in the art described herein. If the molecular beacons probe is added to the reaction mixture prior to 20 amplification it may be possible to monitor the formation of the NASBA product in real-time (Leone et al., Nucleic Acids Research, 1998, Vol 26, 2150-2155).

In a further approach, the molecular beacons technology may be incorporated into the primer 2 oligonucleotide allowing real-time monitoring of the NASBA reaction without the need for a separate hybridisation probe.

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In a still further approach the products of the NASBA reaction may be monitored using a generic labelled detection probe which hybridises to a nucleotide sequence in the 5' terminus of the primer 2 oligonucleotide. This is equivalent to the "NucliSens" detection system supplied by Organon Teknika. In this system specificity for NASBA

products derived from the target HPV mRNA may be conferred by using HPV-specific capture probes comprising probe oligonucleotides as described herein attached to a solid support such as a magnetic microbead. Most preferably the generic labelled detection probe is the ECLTM detection probe supplied by Organon Teknika. NASBA amplicons are hybridized to the HPV-specific capture probes and the generic ECL probe (via a complementary sequence on primer 2).

Following hybridization the bead/amplicon/ECL probe complexes may be captured at the magnet electrode of an automatic ECL reader (e.g. the NucliSens reader supplied by Organon Teknika. Subsequently, a voltage pulse triggers the ECL reaction.

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Also provided by the invention are reagent kits for use in the detection of HPV by NASBA, the kits comprising a primer-pair cocktail according to the The reagent kits may further comprise a mixture of enzymes required for the NASBA reaction, 20 specifically an enzyme mixture containing an RNA directed DNA polymerase (e.g. a reverse transcriptase), a ribonuclease that hydrolyses the RNA strand of an RNA-DNA hybrid without hydrolysing single or double stranded RNA or DNA (e.g. RNaseH) and an RNA 25 The RNA polymerase should be one which polymerase. recognises the promoter sequence present in the 5' terminal region of the NASBA P1 primer oligonucleotides in the oligonucleotide primer sets supplied in the reagent kit. The kit may also 30 comprise a supply of NASBA buffer containing the ribonucleosides and deoxyribonucleosides required for The composition of a standard RNA and DNA synthesis. NASBA reaction buffer will be well known to those skilled in the art. 35

In certain embodiments the kit may further contain one or more capture probes, comprising a probe oligonucleotide attached to a solid support as described above, for immobilising the products of a The kit may still further specific NASBA reaction. contain labelled generic detection probes. Advantageously, the detection probes may comprise a sequence of nucleotides complementary to a non-HPV sequence present at the 5' terminal end of the NASBA P2 primer oligonucleotides present in the reagent kit.

In still further embodiments the kit may further contain one or more molecular beacon probes according to the invention. The molecular beacon probes may be supplied as a separate reagent within the kit. Alternatively, the NASBA primers and molecular beacons probe may be supplied as a primer/probe mixture. a mixture including the NASBA P1 and P2 primers and . also a molecular beacons probe is convenient for use in "real-time" NASBA, wherein the NASBA amplification reaction and detection of an amplification product are performed simultaneously in a single reaction vessel.

The invention will be further understood with reference to the following, non-limiting, Example:

Example 1-Real-time NASBA

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Collection and preparation of clinical samples

Cervical cytobrush samples are collected in 9 ml lysis buffer (5M Guanidine thiocyanate) prior to RNA/DNA extraction. Since RNA is best protected in the 5M guanidine thiocyanate at -70°C only 1 ml of the total volume of sample is used for each extraction 2-3 tubes with the RNA/DNA are stored at -167°C and the rest stored at -70°C.

RNA and DNA were automatically isolated according to the "Booms" isolation method from Organon Teknika (Organon Teknika B.V., Boselind 15, P.O. Box 84, 5280 AB Baxtel, The Netherlands; now Biomérieux, 69280 Marcy 1'Etoile, France).

The following procedure was carried out using reagents from the Nuclisens™ Basic Kit, supplied by Organon Teknika. Procedure for n=10 samples:-

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1. Prepare enzyme solution.

Add 55 μ l of enzyme diluent (from Nuclisens Basic Kit; contains sorbitol in aqueous solution) to each of 3 lyophilized enzyme spheres (from Nuclisens Basic

- Kit; contains AMV-RT, RNase H, T7 RNA polymerase and BSA). Leave this enzyme solution at least for 20 minutes at room temperature. Gather the enzyme solutions in one tube, mix well by flicking the tube with your finger, spin down briefly and use within 1
- hour. Final concentrations in the enzyme mix are 375 mM sorbitol, 2.5 μg BSA, 0.08 U RNase H, 32 U T7 RNA polymerase and 6.4 U AMV-reverse transcriptase.
 - 2. Prepare reagent sphere/KCl solution.
- For 10 samples: add 80 μl reagent sphere diluent (from Nuclisens™ Basic Kit; contains Tris/HCl (pH 8.5), 45% DMSO) to the lyophilized reagent sphere (from Nuclisens™ Basic Kit; contains nucleotides, dithiotreitol and MgCl₂) and immediately vortex well.
- 30 Do this with 3 reagent spheres and mix the solutions in one tube.

Add 3 μl NASBA water (from Nuclisens Basic Kit) to the reconstituted reagent sphere solution and mix well.

Add 56 µl of KCl stock solution (from Nuclisens™ Basic Kit) and mix well. Use of this KCl/water mixture will result in NASBA reactions with a final KCl concentration of 70 mM. Final concentrations in the reagent/KCl solution are 1 mM of each dNTP, 2 mM of ATP, UTP and CTP, 1.5 mM GTP, and 0.5 mM ITP, 0.5 mM dithiotreitol, 70 mM KCl, 12 mM MgCl₂, 40 mM Tris-HCl (pH 8.5).

- 3. Prepare primer/probe solution containing target-specific primers and molecular beacon probe. For each target reaction transfer 91 μl of the reagent sphere/KCl solution (prepared in step 2) into a fresh tube. Add 25 μl of primers/molecular beacon probe solution (to give final concentration of ~0.1-0.5 μM each of the sense and antisense primers and ~ 15-70 pmol molecular beacon probe per reaction). Mix well by vortexing. Do not centrifuge.
 - In case less than 10 target RNA amplifications are being performed refer to the table below for the appropriate amounts of reagent sphere solution, KCl/water solution and primers to be used. Primer solutions should be used within 30 minutes after preparation.

Reactions (n)	Reagent sphere	KCI/water (µI)	Primer mix (µl)
	solution (μl)	30	10
10	80	27	9
9	72		8
8	64	24	7
7	56	21	6
<u>/</u>	48	18	
6	40	15	5
5	32	12	4
4		9	3
3		6	2
2	16	3	1
1	8	_15	

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4. Addition of samples For each target RNA reaction: In a 96 well microtiter plate pipette 10 μ l of the primer/probe solution (prepared in step 3) into each of 10 wells. Add 5 μ l nucleic acid extract to each well. Incubate the microtiter plate for 4 minutes at 65 ± 1 °C. Cool to at 41 ± 0.5 °C for 4 minutes. Then to each well add 5 μ l enzyme solution. Immediately place the microtiter plate in a

O fluorescent detection instrument (e.g. Nuclisens Easy Q Analyzer) and start the amplification.

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- 1. An oligonucleotide molecule for use in the detection of mRNA transcribed from the E6 gene of a human papillomavirus, the oligonucleotide comprising any one of sequence numbers 1-133.
 - 2. An oligonucleotide primer for use in the detection of mRNA transcribed from the E6 gene of a human papillomavirus, the oligonucleotide primer being selected from:
- (i) a NASBA P1 primer comprising one of sequence numbers 2, 4, 8, 11, 14, 17, 20, 22, 25, 28, 31, 34, 37, 39, 42, 45, 48, 50, 52, 57, 60, 63, 65, 69, 71, 75, 78, 81, 84 87, 90, 92, 96, 98, 100, 105, 107, 111, 113, 115, 121, 126, 127, 128 Or 129;
- (ii) a NASBA P2 primer comprising one of sequence numbers 1, 3, 7, 10, 13, 16, 19, 21, 24, 27, 30, 33, 36, 38, 41, 44, 47, 49, 51, 56, 59, 62, 64, 68, 70, 74, 77, 80, 83, 86, 89, 91, 95, 97, 99, 104, 106, 110, 112, 114, 120, 103, 131, 132 or 133;
- (iii) a PCR primer comprising one of sequence numbers

 1, 3, 7, 10, 13, 16, 19, 21, 24, 27, 30, 33, 36, 38,

 41, 44, 47, 49, 51, 56, 59, 62, 64, 68, 70, 74, 77,

 80, 83, 86, 89, 91, 95, 97, 99, 104, 106, 110, 112,

 80, 83, 86, 89, 91, 95, 97, 99, 104, 106, 110, 112,

 114, 120, 2, 4, 8, 11, 14, 17, 20, 22, 25, 28, 31, 34,

 114, 120, 2, 4, 8, 50, 52, 57, 60, 63, 65, 69, 71,

 37, 39, 42, 45, 48, 50, 52, 57, 60, 63, 65, 69, 71,

 37, 78, 81, 84, 87, 90, 92, 96, 98, 100, 105, 107, 111,

 113, 115, 121, 126, 127, 128, 129, 130, 131, 132 or

 133.
 - 3. An oligonucleotide primer according to claim
 2 which is a NASBA P1 primer having the sequence
 AATTCTAATACGACTCACTATAGGGAGAAGG-SEQ, wherein SEQ
 represents any one of sequence numbers 2, 4, 8, 11,

14, 17, 20, 22, 25, 28, 31, 34, 37, 39, 42, 45, 48, 50, 52, 57, 60, 63, 65, 69, 71, 75, 78, 81, 84 87, 90, 92, 96, 98, 100, 105, 107, 111, 113, 115, 121, 126, 127, 128 or 129.

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- 4. An oligonucleotide primer according to claim 2 which is a NASBA P2 primer having the sequence GATGCAAGGTCGCATATGAG-SEQ wherein SEQ represents any one of sequence numbers 1, 3, 7, 10, 13, 16, 19, 21,
- 24, 27, 30, 33, 36, 38, 41, 44, 47, 49, 51, 56, 59, 62, 64, 68, 70, 74, 77, 80, 83, 86, 89, 91, 95, 97, 99, 104, 106, 110, 112, 114, 120, 130, 131, 132 or 133.
- 5. An oligonucleotide probe for use in the detection of mRNA transcribed from the E6 gene of a human papillomavirus comprising one of sequence numbers: 5, 6, 9, 12, 15, 18, 23, 26, 29, 32, 35, 40, 43, 46, 53, 54, 55, 58, 61, 66, 67, 72, 73, 76, 82, 85, 88, 93, 94, 101, 102, 103, 108, 109, 116, 117, 118, 119, 122, 130, 131, 132 or 133.
- 6. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 16, comprising one of the following combinations:

an oligonucleotide primer comprising sequence number 1 and an oligonucleotide primer comprising sequence number 2;

an oligonucleotide primer comprising sequence number 3 and an oligonucleotide primer comprising sequence number 4; an oligonucleotide primer comprising sequence number 7 and an oligonucleotide primer comprising sequence number 8;

an oligonucleotide primer comprising sequence number 10 and an oligonucleotide primer comprising sequence number 11:

an oligonucleotide primer comprising one of sequence numbers 126, 127, 128 or 129 and an oligonucleotide primer comprising sequence number 1 or sequence number 3; or

an oligonucleotide primer comprising sequence number 2 or sequence number 4 and an oligonucleotide primer comprising one of sequence numbers 130, 131, 132 or 133.

7. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 18, comprising one of the following combinations:

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an oligonucleotide primer comprising sequence number 13 and an oligonucleotide primer comprising sequence number 14;

number 14;
20 an oligonucleotide primer comprising sequence number
16 and an oligonucleotide primer comprising sequence
number 17;

an oligonucleotide primer comprising sequence number 19 and an oligonucleotide primer comprising sequence number 20; or

number 20; or an oligonucleotide primer comprising sequence number 21 and an oligonucleotide primer comprising sequence number 22.

30 8. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 31, comprising one of the following combinations:

an oligonucleotide primer comprising sequence number 24 and an oligonucleotide primer comprising sequence number 25;

an oligonucleotide primer comprising sequence number 27 and an oligonucleotide primer comprising sequence number 28;

an oligonucleotide primer comprising sequence number 30 and an oligonucleotide primer comprising sequence number 31;

an oligonucleotide primer comprising sequence number 33 and an oligonucleotide primer comprising sequence number 34; or

an oligonucleotide primer comprising sequence number 36 and an oligonucleotide primer comprising sequence number 37.

9. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 33, comprising one of the following combinations:

an oligonucleotide primer comprising sequence number 38 and an oligonucleotide primer comprising sequence number 39;

an oligonucleotide primer comprising sequence number 41 and an oligonucleotide primer comprising sequence number 42; or

an oligonucleotide primer comprising sequence number 44 and an oligonucleotide primer comprising sequence number 45.

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10. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 35, comprising one of the following combinations:

an oligonucleotide primer comprising sequence number 47 and an oligonucleotide primer comprising sequence number 48;

an oligonucleotide primer comprising sequence number 49 and an oligonucleotide primer comprising sequence number 50; or

an oligonucleotide primer comprising sequence number 51 and an oligonucleotide primer comprising sequence number 52.

- 11. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 52, comprising one of the following combinations:
- an oligonucleotide primer comprising sequence number 56 and an oligonucleotide primer comprising sequence number 57; or an oligonucleotide primer comprising sequence number 59 and an oligonucleotide primer comprising sequence number 60.
- 12. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 58, comprising one of the following combinations:
- an oligonucleotide primer comprising sequence number 62 and an oligonucleotide primer comprising sequence number 63;
 an oligonucleotide primer comprising sequence number 64 and an oligonucleotide primer comprising sequence number 65.
 - 13. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 51, comprising one of the following combinations:
 - an oligonucleotide primer comprising sequence number 104 and an oligonucleotide primer comprising sequence number 105; or
 - an oligonucleotide primer comprising sequence number 106 and an oligonucleotide primer comprising sequence number 107.

- 14. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 56, comprising one of the following combinations:
- an oligonucleotide primer comprising sequence number 110 and an oligonucleotide primer comprising sequence number 111;

an oligonucleotide primer comprising sequence number 112 and an oligonucleotide primer comprising sequence

10 number 113;

an oligonucleotide primer comprising sequence number 114 and an oligonucleotide primer comprising sequence number 115;

an oligonucleotide primer comprising sequence number 120 and an oligonucleotide primer comprising sequence number 121.

- 15. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 39, comprising one of the following combinations:
 - an oligonucleotide primer comprising sequence number 80 and an oligonucleotide primer comprising sequence number 81;
- 25 an oligonucleotide primer comprising sequence number 83 and an oligonucleotide primer comprising sequence number 84; or
 - an oligonucleotide primer comprising sequence number 86 and an oligonucleotide primer comprising sequence number 87.
 - 16. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of HPV 45, comprising one of the following combinations:

an oligonucleotide primer comprising sequence number 89 and an oligonucleotide primer comprising sequence number 90;

an oligonucleotide primer comprising sequence number 91 and an oligonucleotide primer comprising sequence number 92;

an oligonucleotide primer comprising sequence number 95 and an oligonucleotide primer comprising sequence number 96;

an oligonucleotide primer comprising sequence number 97 and an oligonucleotide primer comprising sequence number 98; or

an oligonucleotide primer comprising sequence number 99 and an oligonucleotide primer comprising sequence number 100.

17. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of group B HPV, comprising one of the following combinations:

an oligonucleotide primer comprising sequence number 68 and an oligonucleotide primer comprising sequence number 69;

an oligonucleotide primer comprising sequence number 70 and an oligonucleotide primer comprising sequence number 71; or

an oligonucleotide primer comprising sequence number 74 and an oligonucleotide primer comprising sequence number 75.

18. An oligonucleotide primer-pair for use in the detection of mRNA transcripts from the E6 gene of group C HPV, comprising the following combination:

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- 52 an oligonucleotide primer comprising sequence number 77 and an oligonucleotide primer comprising sequence number 78. An oligonucleotide primer-pair according to 5 any one of claims 6 to 18 which comprises a NASBA P1 primer and a NASBA P2 primer. A primer-pair according to claim 19 wherein the NASBA Pl primer includes the sequence 10 AATTCTAATACGACTCACTATAGGGAGAAGG at the 5' end. A primer/probe set comprising a primer-pair according to any one of claims 6 to 20 and at least one oligonucleotide probe specific for amplification 15 products generated using the primer-pair. A method of detecting HPV mRNA in a test sample suspected of containing HPV which comprises performing an amplification reaction on a preparation 20 of nucleic acid isolated from the test sample to amplify a portion of the mRNA transcribed from the E6 gene of HPV, wherein the amplification reaction is performed using a primer-pair according to any one of 25 claims 6 to 18. A method according to claim 22 which comprises performing RT-PCR to amplify a portion of the mRNA transcribed from the E6 gene of HPV. 30 A method according to claim 126 which comprises performing NASBA to amplify a portion of the mRNA transcribed from the E6 gene of HPV . method according to claim 24 which 35 25. comprises:

- 53 -(a) assembling a reaction mixture comprising a primer set as defined in any one of claims 6 to 18, an RNA directed DNA polymerase, a ribonuclease that hydrolyses the RNA strand of an RNA-DNA hybrid without hydrolysing single or double stranded RNA or DNA, an RNA polymerase that recognises said promoter, and . . . 5 ribonucleoside and deoxyribonucleoside triphosphates; incubating said reaction mixture with a preparation of nucleic acid isolated from a test sample suspected of containing HPV under reaction conditions which permit a NASBA amplification 10 reaction; and detecting and/or quantitatively measuring any :HPV-specific product of the NASBA amplification reaction. 15 A method according to claim 25 wherein step (c) comprises real-time detection of an HPV-specific product of the NASBA amplification reaction. A method according to claim 25 or claim 26 20 wherein the reaction mixture further comprises a molecular beacons probe oligonucleotide and the formation of any HPV-specific NASBA product in the NASBA reaction is monitored by detecting fluorescence from the fluorescent moiety included in the molecular 25 beacons probe. A method according to claim 25 or claim 26 which comprises the further step of capturing the NASBA reaction product by hybridisation to a probe 30 oligonucleotide attached to a solid support. A reagent kit for use in the detection of HPV by NASBA, the kit comprising an oligonucleotide primer-pair as defined in claim 19 and optionally an 35 enzyme mixture comprising an RNA directed DNA

polymerase, a ribonuclease that hydrolyses the RNA strand of an RNA-DNA hybrid without hydrolysing single or double stranded RNA or DNA, and an RNA polymerase that recognises the promoter sequence present in at least one NASBA P1 primer oligonucleotide included in the reagent kit.

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